

STIC Search Report

STIC Database Tracking when her

TO: Angela Martin Location: REM 6B61

Art Unit: 1745 January 4, 2006

Case Serial Number: 10/600452

From: Mei Huang Location: EIC 1700

REMSEN 4B28

Phone: 571/272-3952 Mei.huang@uspto.gov

Search Notes

Examiner Martin,

- 29 answers (L27) shown on page 3-73.

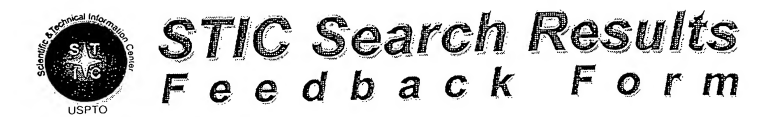
- 27 answers (L28) were retrieved using the secondary keywords "nonaq." etc. (see page 73-75) and were displayed only by titles except answer 24 of 27.

If you have any questions or if you would like to refine the search query, please feel free to contact me.

Thank you for using STIC services!

Mei Huang





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Questions about the scope or the results of the search? Contact the EIC searcher or contact:

Kathleen Fuller, EIC 1700 Team Leader 571/272-2505 REMSEN 4B28

Voluntary Results Feedback Form
 I am an examiner in Workgroup: Example: 1713 Relevant prior art found, search results used as follows:
102 rejection
103 rejection
Cited as being of interest.
Helped examiner better understand the invention.
Helped examiner better understand the state of the art in their technology.
Types of relevant prior art found:
Foreign Patent(s)
 Non-Patent Literature (journal articles, conference proceedings, new product announcements etc.)
> Relevant prior art not found:
Results verified the lack of relevant prior art (helped determine patentability).
Results were not useful in determining patentability or understanding the invention.
Comments:

Server and the Alexander Server Serve

SEARCH REQUEST FORM

Scientific and Technical Information Center

Requester's Full Name: Art Unit: 1745 Phone Numb Mail Box and Bldg/Room Location: Re	Maria E ber 30 <u>571-273-12</u> a 6 <u>361</u> Results	Examiner #: 76027 Date: 1/3/05 & Serial Number: 10/600,452 s Format Preferred (circle): PAPER DISK E-MAIL
If more than one search is submitted		searches in order of need.
'Include the elected species or structures, keywo	ords, synonyms, acronym may have a special mean	specifically as possible the subject matter to be searched. ns, and registry numbers, and combine with the concept or ning. Give examples or relevant citations, authors, etc, if ostract.
		lutron + Nonaqueous secondary battery
Inventors (please provide full names): 2	Marked	
Divini Pir Pir (/a-	1-2 : 0	7/25/47
Earliest Priority Filing Date: $\frac{6/23}{2}$	•	
appropriate serial number.		rent, child, divisional, or issued patent numbers) along with the
Monaqueous electroly	, te Celectr.	slytic solution)
Formula I in cla	in I when	rein X is a halogen (Bronie (Br)),
Chlorine (Cl), Fluo	rine (F), }	Todine (I)
		SCIENTIFIC REFERENCE BR Sci 2 rech Inf - Cn ^t
		JAN 0 3 Kitho
		Pat. & T.M. Office
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	•	
*********	************	*********
$\{V(\lambda)\}$	ype of Search	Vendors and cost where applicable
:	A Sequence (#)	STN
	A Sequence (#)	Dialog
	tructure (#)	Questel/Orbit
1/4/01	ibliographic	Lexis/Nexis
20	itigation	Sequence Systems
	atent Family	WWW/Internet
1.2	ther	Other (specify)

PTO-1590 (8-01)

Other ·

Online Time: _

AMENDMENTS TO THE CLAIMS:

This listing of claims will replace all prior versions, and listings, of claims in the application:

LISTING OF CLAIMS:

1. (original) A nonaqueous electrolytic solution comprising an electrolyte salt dissolved in an organic solvent, which contains a silicon compound represented by formula (I):

$$R_1 - \begin{cases} R_2 \\ S_1 \\ R_3 \end{cases}$$
 (1)

wherein R_1 represents an alkenyl group having 2 to 10 carbon atoms; R_2 and R_3 each represent an alkyl group having 1 to 10 carbon atoms, an alkoxy group having 1 to 10 carbon atoms, an alkenyl group having 2 to 10 carbon atoms or a halogen atom; and X represents a halogen atom.

- 2. (original) The nonaqueous electrolytic solution according to claim 1, wherein R_1 is a vinyl group.
- 3. (original) The nonaqueous electrolytic solution according to claim 1, wherein at least one of R_2 and R_3 is a methyl group.
- 4. (original) The nonaqueous electrolytic solution caccording to claim 1, wherein X is a fluorine atom.

5. (original) The nonaqueous electrolytic solution according to claim 1, wherein R_1 is a vinyl group, R_2 and R_3 are each a methyl group, and X is a fluorine atom.

- 6. (original) The nonaqueous electrolytic solution according to claim 1, wherein the organic solvent contains at least one member selected from the group consisting of a cyclic carbonate compound, a cyclic ester compound, a sulfone compound, a sulfoxide compound, an amide compound, an acyclic carbonate compound, an acyclic ether compound, a cyclic ether compound, and an acyclic ester compound.
- 7. (original) The nonaqueous electrolytic solution according to claim 1, wherein the organic solvent contains at least one cyclic carbonate compound and at least one acyclic carbonate compound.
- 8. (original) The nonaqueous electrolytic solution according to claim 7, wherein the cyclic carbonate compound comprises ethylene carbonate.
- 9. (original) The nonaqueous electrolytic solution according to claim 7, wherein the cyclic carbonate compound comprises ethylene carbonate and 1,2-butylene carbonate.
- 10. (original) The nonaqueous electrolytic solution according to claim 7, wherein the acyclic carbonate compound comprises at least one member selected from the group consisting of dimethyl carbonate, ethylmethyl carbonate, and diethyl carbonate.

11. (original) The nonaqueous electrolytic solution according to claim 7, which further contains a carboxylic ester compound represented by formula (II):

$$CnH_{2n+1}$$
 $C - 0 - R$ (II)

wherein R represents an alkyl group having 1 to 4 carbon atoms; and n represents 0, 1 or 2.

- 12. (original) The nonaqueous electrolytic solution according to claim 1, wherein the electrolyte salt is at least one member selected from the group consisting of LiPF₆, LiBF₄, LiClO₄, LiAsF₆, LiCF₃SO₃, LiN(CF₃SO₂)₂, LiC(CF₃SO₂)₃, an LiCF₃SO₃ derivative, an LiN(CF₃SO₂)₂ derivative, and an LiC(CF₃SO₂)₃ derivative.
- 13. (original) The nonaqueous electrolytic solution according to claim 1, wherein the silicon compound represented by formula (I) is present in an amount of 0.05 to 5% by volume.
- 14. (original) A nonaqueous secondary battery comprising the nonaqueous electrolytic solution according to claim 1.
- 15. (new) The nonaqueous electrolytic solution according to claim 1, further comprising at least 5% by weight of a flame retardant, based on the total organic solvent.

16. (new) The nonaqueous electrolytic solution according to claim 15, wherein the flame retardant is one of a halogen and a phosphorus.

- 17. (new) The nonaqueous electrolytic solution according to claim 16, wherein the flame retardant is a phosphoric ester.
- 18. (new) The nonaqueous electrolytic solution according to claim 1, further comprising 10% to 50% by weight of a flame retardant, based on the total organic solvent.
- 19. (new) The nonaqueous electrolytic solution according to claim 18, wherein the flame retardant is one of a halogen and a phosphorus.
- 20. (new) The nonaqueous electrolytic solution according to claim 19, wherein the flame retardant is a phosphoric ester.

SAV L12 ANG452B/A D SAV

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FILE 'HCAPLUS' ENTERED AT 15:27:23 ON 04 JAN 2006
           3434 SEA L12
L14
L15
         461850 SEA ELECTROLY?
L16
             25 SEA L14 AND L15
              1 SEA L1 AND L16
L17
L18
         227492 SEA BATTERY OR BATTERIES OR (ELECTROCHEM? OR ELECTROLY?
                OR GALVAN? OR WET OR DRY OR PRIMARY OR SECONDARY) (2A) (CEL
                L OR CELLS)
             18 SEA L18 AND L14
L19
L20
             29 SEA L16 OR L19
                D L11 QUE STAT
L21
          45715 SEA NONAQ# OR NONAQUEOUS? OR NONWATER? OR NONH2O OR
               'NON(A) (AQ# OR AQUEOUS? OR WATER? OR H2O)
L22
                QUE (52 OR 72)/SC,SX
L23
             22 SEA L14 AND L21
L24
            36 SEA L14 AND L22
L25
            50 SEA L23 OR L24
L26
             8 SEA L23 AND L24
L27
             29 SEA L20 OR L26
L28
             27 SEA (L23 OR L24) NOT L27
     FILE 'REGISTRY' ENTERED AT 15:44:01 ON 04 JAN 2006
L29
              5 SEA L2 AND ?CARBONAT?/CNS
                D SCA
     FILE 'HCAPLUS' ENTERED AT 15:45:33 ON 04 JAN 2006
          13428 SEA L29
L30
L31
             10 SEA L14 AND L30
L32
             33 SEA L31 OR L27
L33
             5 SEA L31 AND (L21 OR L22)
L34
             29 SEA L33 OR L27
L35
             4 SEA L32 NOT L34
                D SCA TI
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FILE 'REGISTRY' ENTERED AT 15:51:48 ON 04 JAN 2006

=> d 19 que stat L9 STR

VAR G1=8/6/X
NODE ATTRIBUTES:
CONNECT IS E1 RC AT 1
CONNECT IS E1 RC AT 7
CONNECT IS E1 RC AT 8
DEFAULT MLEVEL IS ATOM
GGCAT IS UNS AT 1
GGCAT IS SAT AT 7
DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS

STEREO ATTRIBUTES: NONE

=> fil hcap

FILE 'HCAPLUS' ENTERED AT 15:52:29 ON 04 JAN 2006
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
COPYRIGHT (C) 2006 AMERICAN CHEMICAL SOCIETY (ACS)

=> d 127 ibib abs hitstr hitind 1-YOU HAVE REQUESTED DATA FROM 29 ANSWERS - CONTINUE? Y/(N):y

L27 ANSWER 1 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2005:1220765 HCAPLUS

DOCUMENT NUMBER:

143:463159

TITLE:

Composition of nonaqueous

electrolyte solution and secondary

nonaqueous electrolyte

battery thereof

INVENTOR(S): Fukaya, Atsushi; Usami, Kyohei; Awano, Naomi; Tachikawa, Hiroyuki; Taki, Takayuki PATENT ASSIGNEE(S): Asahi Denka Co., Ltd., Japan SOURCE: PCT Int. Appl., 33 pp. CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: Japanese FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE _ _ _ _ WO 2005109561 **A1** 20051117 WO 2005-JP8146 200504 AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, KE, KG, KM, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW,

20051222

GN, GQ, GW, ML, MR, NE, SN, TD, TG

A2

AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA,

200504

15

Α

PRIORITY APPLN. INFO.:

JP 2005353579

JP 2004-140889

JP 2005-117991

200405 11

GΙ

II

$$\begin{array}{c|c}
R^{1} & R^{4} \\
R^{2} - Si & X - Si \\
R^{3} & R^{6}
\end{array}$$

$$\begin{array}{c}
R^{4} \\
R^{5} \\
R^{5}
\end{array}$$

$$\begin{array}{c}
R^{8} \\
R^{7} - Si - X \\
R^{9}
\end{array}$$

The electrolyte soln. contains a mixed org. solvent comprising ethylene carbonate 20-35, Et Me carbonate 35-45, Me2CO3 15-35, and Et2CO3 an/or (C3H7)2CO3 3-15 vo.%. The electrolyte soln. contains inorg. and/or org. Li salt as solute. and may also contain an additive I [R1-6= (ether bonding contg.) alkyl, alkoxy, alkenyl, alkenyloxy, aryl, or aryloxy groups; n = 0-5, , and X = a bond, O, Alkylene, alkylene dioxy, alkenylene, alkenylene dioxy, alkenylene, alkenylene dioxy, arylene, or arylene dioxy groups when n = 1-5, and ≥1 of R1-6 and X has an unsatd. group] or II (R7 = C2-10 alkenyl group, R8 and R9 = C1-10 alkyl, alkoxy or C2-10 alkenyl group or halogen, X = halogen).

IT 1719-58-0 210362-80-4

RL: MOA (Modifier or additive use); USES (Uses) (compns. of org. solvent mixts. for electrolyte solns. in secondary lithium batteries)

RN 1719-58-0 HCAPLUS

CN Silane, chloroethenyldimethyl- (9CI) (CA INDEX NAME)

RN 210362-80-4 HCAPLUS CN Silane, fluorodimethyl-2-propenyl- (9CI) (CA INDEX NAME)

IC ICM H01M010-40

CC **52-2** (Electrochemical, Radiational, and Thermal Energy Technology)

ST secondary lithium battery electrolyte solvent compn

IT Battery electrolytes

(compns. of org. solvent mixts. for electrolyte solns. in secondary lithium batteries)

IT 96-49-1, Ethylene carbonate 105-58-8, Diethyl carbonate 108-32-7, Propylene carbonate 616-38-6, Dimethyl carbonate 623-53-0, Ethyl methyl carbonate 21324-40-3, Lithium hexafluorophosphate

RL: DEV (Device component use); USES (Uses) (compns. of org. solvent mixts. for electrolyte solns. in secondary lithium batteries)

IT 1719-58-0 17955-81-6 18645-49-3 210362-80-4
RL: MOA (Modifier or additive use); USES (Uses)
(compns. of org. solvent mixts. for electrolyte solns. in secondary lithium batteries)

REFERENCE COUNT: 13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE

IN THE RE FORMAT

L27 ANSWER 2 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2005:667917 HCAPLUS

DOCUMENT NUMBER:

143:327229

TITLE:

Novel polymeric systems for lithium ion

batteries gel electrolytes II.

Hybrid cross-linked poly(fluorosilicon-ethylene

oxide)

AUTHOR(S):

Appetecchi, G. B.; Alessandrini, F.; Passerini,

S.; Caporiccio, G.; Boutevin, B.;

Guida-PietraSanta, F.

CORPORATE SOURCE:

Idrocomb C.R. Casaccia, ENEA, Rome, 00060, Italy

SOURCE: Electrochimica Acta (2005), 50(22), 4396-4404

CODEN: ELCAAV; ISSN: 0013-4686

PUBLISHER: Elsevier B.V.

DOCUMENT TYPE:

Journal English

LANGUAGE: Crosslinked, self-supporting, membranes for lithium ion AB battery gel electrolytes were obtained by crosslinking a mixt. of polyfluorosilicone (PFSi) and polysilicone contg. ethylene oxide (EO) units [P(Si-EO)]. The membranes were also reinforced with nanosized silica. The two polymer precursors were synthesized with functional groups capable to form inter-mol. crosslinking, thus obtaining three-dimensional, polymer matrixes. The precursors were dissolved in a common solvent and cross-linked to obtain free-standing PFSi/P(Si-EO):SiO2 composite films. latter were undergone to swelling processes in (nonaq., aprotic, lithium salt contg.) electrolytic solns. to obtain gel-type polymer electrolytes. The properties of the swelled PFSi/P(Si-EO):SiO2 samples were evaluated as a function of the electrolytic solns. and the dipping The PFSi/P(Si-EO):SiO2 membranes exhibited large swelling properties, high ionic cond. and good electrochem. stability. 1719-58-0DP, reaction products with polysiloxanes, polymers IT RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation);

PREP (Properties); RCT (Reactant); SPN (Synthetic preparation);
PREP (Preparation); RACT (Reactant or reagent)

(prepn. and elec. properties of crosslinked poly(fluorosilicon-

ethylene oxide) lithium ion batteries gel electrolyte)

RN 1719-58-0 HCAPLUS

CN Silane, chloroethenyldimethyl- (9CI) (CA INDEX NAME)

IT 1719-58-0, Chlorodimethylvinylsilane

RL: RCT (Reactant); RACT (Reactant or reagent)
(prepn. and elec. properties of crosslinked poly(fluorosiliconethylene oxide) lithium ion batteries gel
electrolyte)

RN 1719-58-0 HCAPLUS

CN Silane, chloroethenyldimethyl- (9CI) (CA INDEX NAME)

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\begin{array}{c} \text{Cl} \\ | \\ \text{Me-Si-CH----} \text{CH}_2 \\ | \\ \text{Me} \end{array}
```

CC 38-3 (Plastics Fabrication and Uses)

Section cross-reference(s): 52

ST lithium ion battery gel electrolyte crosslinked polyfluorosilicon ethylene oxide

IT Secondary batteries

(lithium ion; prepn. and elec. properties of crosslinked poly(fluorosilicon-ethylene oxide) lithium ion batteries gel electrolyte)

IT Polysiloxanes, uses

RL: DEV (Device component use); POF (Polymer in formulation); PRP (Properties); SPN (Synthetic preparation); PREP (Preparation); USES (Uses)

(polyoxyalkylene-, fluorine-contg.; prepn. and elec. properties of crosslinked poly(fluorosilicon-ethylene oxide) lithium ion batteries gel electrolyte)

IT Fluoropolymers, uses

RL: DEV (Device component use); POF (Polymer in formulation); PRP (Properties); SPN (Synthetic preparation); PREP (Preparation); USES (Uses)

(polyoxyalkylene-siloxane-; prepn. and elec. properties of crosslinked poly(fluorosilicon-ethylene oxide) lithium ion batteries gel electrolyte)

IT Electric impedance

Ionic conductivity

Membranes, nonbiological

Polymer electrolytes

Swelling, physical

(prepn. and elec. properties of crosslinked poly(fluorosiliconethylene oxide) lithium ion batteries gel

electrolyte)

IT Polyoxyalkylenes, uses

RL: DEV (Device component use); POF (Polymer in formulation); PRP (Properties); USES (Uses)

(prepn. and elec. properties of crosslinked poly(fluorosiliconethylene oxide) lithium ion batteries gel

electrolyte)

IT Polyoxyalkylenes, uses

RL: DEV (Device component use); POF (Polymer in formulation); PRP (Properties); SPN (Synthetic preparation); PREP (Preparation); USES (Uses)

(siloxane-, fluorine-contg.; prepn. and elec. properties of crosslinked poly(fluorosilicon-ethylene oxide) lithium ion batteries gel electrolyte)

IT 865316-04-7DP, lithium complexes, perchlorate- or hexafluorophosphate-contg. 865316-04-7P

RL: DEV (Device component use); POF (Polymer in formulation); PRP (Properties); SPN (Synthetic preparation); PREP (Preparation); USES (Uses)

(crosslinked; prepn. and elec. properties of crosslinked poly(fluorosilicon-ethylene oxide) lithium ion **batteries** gel **electrolyte**)

IT 96-48-0 96-49-1, 1,3-Dioxolan-2-one 105-58-8 616-38-6, Dimethyl carbonate

RL: NUU (Other use, unclassified); USES (Uses)

(electrolyte soln.-contg.; prepn. and elec. properties of crosslinked poly(fluorosilicon-ethylene oxide) lithium ion batteries gel electrolyte)

IT 7631-86-9, Silica, uses

RL: MOA (Modifier or additive use); USES (Uses)
(nanometric; prepn. and elec. properties of crosslinked poly(fluorosilicon-ethylene oxide) lithium ion batteries gel electrolyte)

IT 25322-68-3, PEO

RL: DEV (Device component use); POF (Polymer in formulation); PRP (Properties); USES (Uses)

(prepn. and elec. properties of crosslinked poly(fluorosiliconethylene oxide) lithium ion **batteries** gel **electrolyte**)

TT 7439-93-2DP, Lithium, crosslinked poly(fluorosilicon-ethylene oxide) complexes, perchlorate- or hexafluorophosphate-contg.

RL: DEV (Device component use); POF (Polymer in formulation); PRP (Properties); SPN (Synthetic preparation); PREP (Preparation); USES (Uses)

(prepn. and elec. properties of crosslinked poly(fluorosiliconethylene oxide) lithium ion **batteries** gel **electrolyte**)

IT 1719-58-0DP, reaction products with polysiloxanes, polymers

865316-00-3DP, dimethylvinylsilyl-terminated 865316-02-5DP, dimethylvinylsilyl-terminated RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (prepn. and elec. properties of crosslinked poly(fluorosiliconethylene oxide) lithium ion batteries gel electrolyte) IT 74-85-1, Ethene, reactions 335-70-6 1066-35-9 **1719-58-0** , Chlorodimethylvinylsilane 58185-54-9, 4,7,10,13,16-Pentaoxanonadeca-1,18-diene RL: RCT (Reactant); RACT (Reactant or reagent) (prepn. and elec. properties of crosslinked poly(fluorosiliconethylene oxide) lithium ion batteries gel electrolyte) IT 35192-49-5P 865316-01-4P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (prepn. and elec. properties of crosslinked poly(fluorosiliconethylene oxide) lithium ion batteries gel electrolyte) REFERENCE COUNT: 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L27 ANSWER 3 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:521417 HCAPLUS

DOCUMENT NUMBER: 143:194818

TITLE: Two new siloxanic proton-conducting membranes

AUTHOR(S): Di Noto, Vito; Vittadello, Michele

CORPORATE SOURCE: Dipartimento di Scienze Chimiche, Universita di

Padova, Padua, I-35135, Italy

SOURCE: Electrochimica Acta (2005), 50(19), 3998-4006

CODEN: ELCAAV; ISSN: 0013-4686

PUBLISHER: Elsevier B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

AB The development of stable polymer electrolytes having good proton cond., low cost and operating at medium temps. represent a crucial step in the evolution of polymer electrolyte fuel cells. We describe two new siloxanic proton-conducting membranes that were synthesized through a two-stage protocol. In the first stage, a poly(Me hydro siloxane) precursor (P) bearing siloxane side chains with sulfonic acid groups was prepd. In the second step, the hydrolysis of pristine precursor or its deriv. obtained by grafting siloxane chains on P yielded two types of

membranes with the formulas $\{Si(CH3)30[Si(CH3)H0]21.26 - [Si(CH3)((CH2)3S03H)0]1.8 - [Si(CH3)((CH2)3Si(CH3)20-)-0]14 - Si(CH3)3\}n$ (A) and $\{Si(CH3)30[Si(CH3)H0]21.26 - [Si(CH3)((CH2)3S03H)0]1.8 - [Si(CH3)((CH2)3(Si(CH3)20-)w)-0]v[Si(CH3)((CH2)3Si(CH3)20-)-0]14-vSi(CH3)3\}n$ (B), with w=20.31. Polymer membranes of A and B were prepd. by means of a hot-pressing process at 80 °C and 10 t/cm2. SEM showed that A and B are rubbery materials with rough and transparent surfaces. Thermogravimetric investigations performed under air atm. disclosed that A and B are thermally stable up to at least 198 °C. DSC measurements yielded T g(s) of -44 and -60 °C for A and B, resp. The polymers exhibit ionic exchange capacities of 0.33 (A) and 0.15 m-eq/g (B). FT-IR and FT-Raman investigations revealed that the polymers consist of reticulated siloxane networks with pendant silicone chains having sulfonic acid groups.

IT 4028-23-3, Allylchlorodimethylsilane

RL: RCT (Reactant); RACT (Reactant or reagent)

(hydrosilylation agent; siloxanic proton-conducting membranes)

RN 4028-23-3 HCAPLUS

CN Silane, chlorodimethyl-2-propenyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \texttt{Cl} \\ | \\ \texttt{Me-Si-CH}_2-\texttt{CH} \end{array} \\ = \begin{array}{c} \texttt{CH}_2 \\ | \\ \texttt{Me} \end{array}$$

CC 38-3 (Plastics Fabrication and Uses) Section cross-reference(s): 35, 37

4028-23-3, Allylchlorodimethylsilane 14418-84-9,

2-Propene-1-sulfonyl chloride

RL: RCT (Reactant); RACT (Reactant or reagent)

24

(hydrosilylation agent; siloxanic proton-conducting membranes)

REFERENCE COUNT:

IT

THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE

IN THE RE FORMAT

L27 ANSWER 4 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:21131 HCAPLUS

DOCUMENT NUMBER: 142:414406

TITLE: Conducting agent for alkaline Zn/MnO2

battery and its production

INVENTOR(S): Su, Lixiao

PATENT ASSIGNEE(S):

Yuanhan Graphite Coating Factory, Shaowu,

Fujian, Peop. Rep. China

SOURCE:

Faming Zhuanli Shenqing Gongkai Shuomingshu, 10

pp.

CODEN: CNXXEV

DOCUMENT TYPE:

Patent Chinese

LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1481038	Α	20040310	CN 2003-149783	
·				200308
				06
PRIORITY APPLN. INFO.:			CN 2003-149783	
				200308
				06

AB The conducting agent with Fe content <20 x 10-6 is composed of 181-377 part resin compn. soln. and 50-100 part conductive powder. The resin compn. soln. is composed of vinyl resin 10- 30, copolymer resin 5-15, coupling agent 1-2, ketone solvent 80- 160, ester solvent 20-40, benzene soln. 20-40, alkane solvent 20-40, THF 20-40, and plasticizer 5-10 part. The vinyl resin is polyvinyl chloride, poly(vinyl butyral), paste polyvinyl chloride resin, poly(vinylidene dichloride) resin, and/or poly(perchloroethylene) resin. copolymer resin is hydroxy-(carboxyl, or ester)- contq. vinyl chloride-vinyl acetate copolymer resin, vinyl chloride-vinyl acetate copolymer resin, and/or vinylidene dichloride-acrylonitrile copolymer resin. The coupling agent is vinyltrichlorosilane, methyltrimethoxysilane, vinyltriethoxysilane, and/or vinyltrimethoxysilane. The plasticizer is diisooctyl phthalate, dioctyl phthalate, di-Bu phthalate, dioctyl adipate, or diisodecyl phthalate. The conductive powder is graphite or Carbon black.

IT 75-94-5, Vinyltrichlorosilane

RL: DEV (Device component use); USES (Uses) (coupling agent; conducting agent for alk. Zn/MnO2 battery and its prodn.)

RN 75-94-5 HCAPLUS

CN Silane, trichloroethenyl- (9CI) (CA INDEX NAME)

```
Cl_3Si - CH = CH_2
IC
     ICM H01M002-20
     ICS H01M006-02; H01B001-24; C09D005-24; C09J009-02
CC
     52-2 (Electrochemical, Radiational, and Thermal Energy Technology)
ST
     conducting agent alk zinc manganese dioxide battery
IT
     Esters, uses
   . RL: DEV (Device component use); USES (Uses)
        (arom.; conducting agent for alk. Zn/MnO2 battery and
        its prodn.)
IT
     Polymers, uses
     RL: DEV (Device component use); POF (Polymer in formulation); USES
        (co-; conducting agent for alk. Zn/MnO2 battery and its
        prodn.)
     Coupling agents
IT
     Plasticizers
     Solvents
        (conducting agent for alk. Zn/MnO2 battery and its
        prodn.)
ΙT
     Alkanes, uses
     Ketones, uses
     Silanes
     RL: DEV (Device component use); USES (Uses)
        (conducting agent for alk. Zn/MnO2 battery and its
        prodn.)
IT
     Polyvinyl butyrals
     RL: DEV (Device component use); POF (Polymer in formulation); USES
        (conducting agent for alk. Zn/MnO2 battery and its
        prodn.)
IT
     Battery electrodes
        (conductive materials for; conducting agent for alk. Zn/MnO2
        battery and its prodn.)
IT
     Carbon black, uses
     RL: DEV (Device component use); TEM (Technical or engineered
     material use); USES (Uses)
        (conductor; conducting agent for alk. Zn/MnO2 battery
        and its prodn.)
IT
     Vinyl compounds, uses
     RL: DEV (Device component use); POF (Polymer in formulation); USES
     (Uses)
        (polymers; conducting agent for alk. Zn/MnO2 battery
```

and its prodn.) IT 71-43-2, Benzene, uses 109-99-9, Tetrahydrofuran, uses RL: DEV (Device component use); USES (Uses) (conducting agent for alk. Zn/MnO2 battery and its prodn.) IT 9002-85-1, Poly(vinylidene dichloride) 9002-86-2, Polyvinyl chloride 9003-22-9, Vinyl acetate-vinyl chloride copolymer 9003-22-9D, Vinyl acetate-vinyl chloride copolymer, partially hydrolyzed 9010-76-8, Acrylonitrile-vinylidene dichloride copolymer 25135-99-3, Poly(tetrachloroethylene) RL: DEV (Device component use); POF (Polymer in formulation); USES (Uses) (conducting agent for alk. Zn/MnO2 battery and its prodn.) IT 7782-42-5, Graphite, uses RL: DEV (Device component use); TEM (Technical or engineered material use); USES (Uses) (conductor; conducting agent for alk. Zn/MnO2 battery and its prodn.) **75-94-5**, Vinyltrichlorosilane 78-08-0, IT Vinyltriethoxysilane 1185-55-3, Methyltrimethoxysilane 2768-02-7, Vinyltrimethoxysilane RL: DEV (Device component use); USES (Uses) (coupling agent; conducting agent for alk. Zn/MnO2 battery and its prodn.) ΙT 1313-13-9, Manganese dioxide, uses RL: DEV (Device component use); USES (Uses) (electrode material; conducting agent for alk. Zn/MnO2 battery and its prodn.) IT7440-66-6, Zinc, uses RL: DEV (Device component use); TEM (Technical or engineered material use); USES (Uses) (electrode material; conducting agent for alk. Zn/MnO2 battery and its prodn.) IT 14280-30-9, Hydroxide, uses RL: DEV (Device component use); USES (Uses) (electrolyte solns.; conducting agent for alk. Zn/MnO2 battery and its prodn.) IT 84-74-2, Dibutyl phthalate 103-23-1, Dioctyl adipate 117-81-7, Diisooctyl phthalate 117-84-0, Di-n-octyl phthalate 26761-40-0,

(plasticizer; conducting agent for alk. Zn/MnO2 battery

RL: DEV (Device component use); USES (Uses)

Diisodecyl phthalate

and its prodn.)

L27 ANSWER 5 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2004:850384 HCAPLUS DOCUMENT NUMBER: 142:41337 TITLE: Novel polymeric systems for lithium-ion batteries gel electrolytes I. Cross-linked polyfluorosilicone AUTHOR(S): Appetecchi, G. B.; Alessandrini, F.; Passerini, S.; Caporiccio, G.; Boutevin, B.; Guida-Pietrasanta, F. Idrocomb C.R. Casaccia, ENEA, Rome, 00060, Italy CORPORATE SOURCE: SOURCE: Electrochimica Acta (2004), 50(1), 149-158 CODEN: ELCAAV; ISSN: 0013-4686 PUBLISHER: Elsevier B.V. DOCUMENT TYPE: Journal LANGUAGE: English AB The study of chem. cross-linked, self-supporting gel-type electrolyte membranes, based on hybrid polyfluorosilicone polymers reinforced with nanosized silica, for lithium-ion battery systems is reported. The polyfluorosilicone materials were selected from their high chem. and thermal stabilities. The precursors were synthesized with functional groups capable to form inter-mol. crosslinking, thus obtaining three-dimensional polymer matrixes. The latter were undergone to swelling processes in (nonaq., lithium salt contg.) electrolytic solns. to obtain gel-type polymer electrolytes. Several kinds of membranes, based on different types of polyfluorosilicone precursor, were prepd. and characterized in terms of swelling behavior, ionic cond. and electrochem. stability. The properties of the swelled matrixes were evaluated as a function of dipping time, temp., kind of electrolytic soln. and crosslinking initiator content. IT 124-70-9DP, Dichloromethylvinylsilane, hydrolyzed, copolymers with fluoroalkyl methylsiloxy- and dimethylsiloxy- group contg. telomers 1719-58-0DP, Chlorodimethylvinylsilane, hydrolyzed, copolymers with fluoroalkyl methylsiloxy- and dimethylsiloxy- group contg. telomers RL: PEP (Physical, engineering or chemical process); PRP (Properties); PYP (Physical process); SPN (Synthetic preparation); PREP (Preparation); PROC (Process) (composites with silica; cross-linked polyfluorosilicons as novel

RN 124-70-9 HCAPLUS

electrolytes)

CN Silane, dichloroethenylmethyl- (9CI) (CA INDEX NAME)

polymeric systems for lithium-ion batteries gel

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Cl
Me-Si-CH=CH_2
    C1
RN
     1719-58-0 HCAPLUS
     Silane, chloroethenyldimethyl- (9CI) (CA INDEX NAME)
CN
   Cl
Me-Si-CH-CH2
   Me
     52-2 (Electrochemical, Radiational, and Thermal Energy
CC
     Technology)
     Section cross-reference(s): 38, 39, 49, 76
     polymer gel electrolyte lithium ion battery
st
     silica composite polyfluorosilicone
IT
     Battery electrolytes
     Gels
     Polymer electrolytes
     Swelling, physical
        (cross-linked polyfluorosilicons as novel polymeric systems for
        lithium-ion batteries gel electrolytes)
IT
     Fluoropolymers, preparation
     RL: PEP (Physical, engineering or chemical process); PRP
     (Properties); PYP (Physical process); SPN (Synthetic preparation);
     PREP (Preparation); PROC (Process)
        (di-Me siloxane-, fluoroalkyl- and vinyl group-contg., vinyl
        group-terminated; cross-linked polyfluorosilicons as novel
        polymeric systems for lithium-ion batteries gel
        electrolytes)
IT
     Polysiloxanes, preparation
     RL: PEP (Physical, engineering or chemical process); PRP
     (Properties); PYP (Physical process); SPN (Synthetic preparation);
     PREP (Preparation); PROC (Process)
        (di-Me, fluoroalkyl- and vinyl group-contg., vinyl
        group-terminated; cross-linked polyfluorosilicons as novel
```

polymeric systems for lithium-ion batteries gel

```
electrolytes)
IT
     Silicone rubber, preparation
     RL: PEP (Physical, engineering or chemical process); PRP
     (Properties); PYP (Physical process); SPN (Synthetic preparation);
     PREP (Preparation); PROC (Process)
        (fluorine-contg.; cross-linked polyfluorosilicons as novel
        polymeric systems for lithium-ion batteries gel
        electrolytes)
IT
     Telomers (polymers)
     RL: PEP (Physical, engineering or chemical process); PRP
     (Properties); PYP (Physical process); SPN (Synthetic preparation);
     PREP (Preparation); PROC (Process)
        (fluoropolymers, copolymers with dichloromethyl and
        chlorodimethylsilane derivs., composites with silica;
        cross-linked polyfluorosilicons as novel polymeric systems for
        lithium-ion batteries gel electrolytes)
IT
     Secondary batteries
        (lithium; cross-linked polyfluorosilicons as novel polymeric
        systems for lithium-ion batteries gel
        electrolytes)
IT
     Ionic conductivity
        (of gel polymer electrolyte composites; cross-linked
        polyfluorosilicons as novel polymeric systems for lithium-ion
        batteries gel electrolytes)
     Electric current-potential relationship
IT
        (of gel polymer electrolytes in electrolytic
        solns.; cross-linked polyfluorosilicons as novel polymeric
        systems for lithium-ion batteries gel
        electrolytes)
IT
     Glass transition temperature
        (of polysiloxanes, di-Me, fluoroalkyl- and vinyl group-contg.
        precursors and crosslinked silica-composite membranes;
        cross-linked polyfluorosilicons as novel polymeric systems for
        lithium-ion batteries gel electrolytes)
IT
     Fluoro rubber
     RL: PEP (Physical, engineering or chemical process); PRP
     (Properties); PYP (Physical process); SPN (Synthetic preparation);
     PREP (Preparation); PROC (Process)
        (silicone; cross-linked polyfluorosilicons as novel polymeric
        systems for lithium-ion batteries gel
        electrolytes)
IT
     Fluoropolymers, preparation
     RL: PEP (Physical, engineering or chemical process); PRP
     (Properties); PYP (Physical process); SPN (Synthetic preparation);
     PREP (Preparation); PROC (Process)
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(telomers, copolymers with dichloromethyl and chlorodimethylsilane derivs., composites with silica; cross-linked polyfluorosilicons as novel polymeric systems for lithium-ion batteries gel electrolytes)

IT 7631-86-9, Aerosil 200, uses

RL: PEP (Physical, engineering or chemical process); PYP (Physical process); TEM (Technical or engineered material use); PROC (Process); USES (Uses)

(colloidal; cross-linked polyfluorosilicons as novel polymeric systems for lithium-ion **batteries** gel

electrolytes)

IT 7791-03-9, Lithium perchlorate 21324-40-3, Lithium hexafluorophosphate

RL: DEV (Device component use); USES (Uses)
 (composite gel polymer electrolytes with carbonates and
 dimethyl-fluoroalkyl-polysiloxanes; cross-linked
 polyfluorosilicons as novel polymeric systems for lithium-ion
 batteries gel electrolytes)

- 124-70-9DP, Dichloromethylvinylsilane, hydrolyzed, copolymers with fluoroalkyl methylsiloxy- and dimethylsiloxy- group contg. telomers 1066-35-9DP, Chlorodimethylsilane, copolymers with fluoroalkyl methylsiloxy- and dimethylsiloxy- group contg. telomers 1719-58-0DP, Chlorodimethylvinylsilane, hydrolyzed, copolymers with fluoroalkyl methylsiloxy- and dimethylsiloxy- group contg. telomers
 - RL: PEP (Physical, engineering or chemical process); PRP (Properties); PYP (Physical process); SPN (Synthetic preparation); PREP (Preparation); PROC (Process)

(composites with silica; cross-linked polyfluorosilicons as novel polymeric systems for lithium-ion batteries gel electrolytes)

- TT 78-63-7, 2,5-Bis(tert-butylperoxy)-2,5-dimethyl-hexane
 RL: CAT (Catalyst use); USES (Uses)
 - (cross-linked polyfluorosilicons as novel polymeric systems for lithium-ion batteries gel electrolytes)
- TT 74-84-0DP, Ethane, copolymers with 1,6-diiodododecafluorohexane, chlorodimethylsilane, hydrolyzed 1,1,3,3,5,5-Hexamethyltrisiloxane, and hydrolyzed dichloromethylvinylsilane and chlorodimethylvinylsilane 335-70-6DP, 1,8-Diiodohexadecafluorooctane, copolymers with ethane, chlorodimethylsilane, hydrolyzed 1,1,3,3,5,5-Hexamethyltrisiloxane, and hydrolyzed dichloromethylvinylsilane and chlorodimethylvinylsilane 375-80-4DP, 1,6-Diiodododecafluorohexane, copolymers with ethane, chlorodimethylsilane, hydrolyzed 1,1,3,3,5,5-Hexamethyltrisiloxane,

and hydrolyzed dichloromethylvinylsilane and chlorodimethylvinylsilane 591-87-7DP, Allyl acetate, copolymers with chlorodimethylsilane, and hydrolyzed dichloromethylvinylsilane and chlorodimethylvinylsilane and diiodo-terminated copolymers of vinylidene difluoride, hexafluoropropene, tetrafluoroethenevinylidene difluoride 1189-93-1DP, 1,1,3,3,5,5-Hexamethyltrisiloxane, hydrolyzed, copolymers with ethane, 1,6-diiodododecafluorohexane, hydrolyzed dichloromethylvinylsilane and chlorodimethylvinylsilane RL: PEP (Physical, engineering or chemical process); PRP (Properties); PYP (Physical process); SPN (Synthetic preparation); PREP (Preparation); PROC (Process)

(cross-linked polyfluorosilicons as novel polymeric systems for lithium-ion batteries gel electrolytes)

IT 96-48-0, γ-Butyrolactone 96-49-1, Ethylene carbonate 105-58-8, Diethylcarbonate 108-32-7, Propylene carbonate 616-38-6, Dimethyl carbonate

RL: DEV (Device component use); USES (Uses)
 (gel composites with carbonates/lithium salts/dimethylfluoroalkyl-polysiloxanes; cross-linked polyfluorosilicons as
novel polymeric systems for lithium-ion batteries gel
electrolytes)

REFERENCE COUNT:

OUNT: 24 THERE ARE 24 CITED REFERENCES AVAILABLE
FOR THIS RECORD. ALL CITATIONS AVAILABLE
IN THE RE FORMAT

L27 ANSWER 6 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2

2004:493203 HCAPLUS

DOCUMENT NUMBER:

141:40696

TITLE:

Nonaqueous electrolyte

solution and secondary lithium battery

which uses the solution

INVENTOR(S):
Hinohara, Akio; Hayashi, Takeshi

PATENT ASSIGNEE(S): Mitsui Chemicals Inc., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 13 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2004171981	A2	20040617	JP 2002-337756	

200211 21

PRIORITY APPLN. INFO.:

JP 2002-337756

200211 21

AB The electrolyte soln. comprises a Li salt and a nonaq. solvent mixt. contg. a fluorosilane compd. of the structure R1R2R3SiF where R1-3 = F or a C1-12 hydrocarbon group; and ≥1 of R1-3 = a C1-12 hydrocarbon group. The battery has a Li-intercalating anode, a Li-intercalating anode, and the above electrolyte soln.

IT 38755-76-9, Fluorovinyl dimethyl silane
RL: MOA (Modifier or additive use); USES (Uses)
(electrolyte solns. contg. fluorosilane compds. in solvents for secondary lithium batteries)

RN 38755-76-9 HCAPLUS

CN Silane, ethenylfluorodimethyl- (9CI) (CA INDEX NAME)

IC ICM H01M010-40

CC **52-2** (Electrochemical, Radiational, and Thermal Energy Technology)

ST secondary lithium battery electrolyte nonaq solvent fluorosilane compd

IT Battery electrolytes

(electrolyte solns. contg. fluorosilane compds. in solvents for secondary lithium batteries)

IT Secondary batteries

(lithium; electrolyte solns. contg. fluorosilane compds. in solvents for secondary lithium batteries)

IT 96-49-1, Ethylene carbonate 623-53-0, Methyl ethyl carbonate 21324-40-3, Lithium hexafluorophosphate

RL: DEV (Device component use); USES (Uses)

(electrolyte solns. contg. fluorosilane compds. in solvents for secondary lithium batteries)

IT 312-40-3, Difluorodiphenyl silane 368-47-8, Phenyl trifluorosilane 379-50-0, Fluorotriphenyl silane 420-56-4, Fluorotrimethyl silane

872-36-6, Vinylene carbonate 4427-96-7, Vinyl ethylene carbonate 4663-29-0 **38755-76-9**, Fluorovinyl dimethyl silane 96164-66-8, Trifluorohexyl silane RL: MOA (Modifier or additive use); USES (Uses) (electrolyte solns. contg. fluorosilane compds. in

L27 ANSWER 7 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

solvents for secondary lithium batteries)

ACCESSION NUMBER: 2004:413007 HCAPLUS

DOCUMENT NUMBER: 140:424654

TITLE: Method to produce graphite/polymer hybrid

composites

INVENTOR(S): Kasseh, Abdeslam; Chaouki, Jamal; Ennajimi,

Elmekki

PATENT ASSIGNEE(S): Can.

SOURCE: PCT Int. Appl., 32 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PATENT	NO.			KIN	D :	DATE			APPL	ICAT	ION :	NO.		D.	ATE	
- V	 10 2004	- :0419	15		A 1		2004	0521		WO 2	003-	CA17	31				
															_	00311 7	
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	-	-	
		CH,	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	
		GB,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	
		KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	
		MX,	MZ,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	
				SL,			TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	
•		VN,	YU,	ZA,	ZM,	ZW											
	RW:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	
				KG,												-	
				ES,												-	
				SK,			ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	
		•	•	SN,	•												
C	'A 2411	443			AA		2004	0507	(CA 2	002-	2411	443				
															2 ·	00211 7	
PRIORI	TY APP	LN.	INFO	. :						CA 2	002-	24114	443	7	A J	•	
				- •										•	_	00211	

AB Hybrid composites with tailored structure were prepd. using a novel synthesis method based on stable free radical polymn. combined with polymn. compounding. Highly filled and well-dispersed filler/polymer composites have been developed using this method. Grafting polymers onto carbon fillers was carried out in bulk polymn., in soln. and in colloidal dispersion. This method allows control of the percentage of grafted polymers, the architecture of grafted polymers, the length of chains, and the polydispersity index. The org. fillers may be graphite, carbon black and the like in the form of flakes, fibers, colloidal suspensions, films or powders. The synthesis process performed by this method gave grafting percentages of polymers and copolymers ranging from 12 to Graphite/Polymer composites produced herein represent new material used in bipolar plates for fuel cells and filter press electrolyzers and as composite material for automobiles and aircraft structures.

IT 17196-12-2, 7-Octenyldimethylchlorosilane RL: RCT (Reactant); RACT (Reactant or reagent)

(free radical graft polymn. in method to produce graphite/polymer hybrid composites)

RN 17196-12-2 HCAPLUS

CN Silane, chlorodimethyl-7-octenyl- (8CI, 9CI) (CA INDEX NAME)

$$^{\text{Cl}}$$
 $^{\text{Me}-\text{Si}-\text{(CH}_2)}_{6}$
 $^{\text{CH}=\text{CH}=\text{CH}_2}$
 $^{\text{Me}}$

IC ICM C08J005-06

ICS C08J005-24; C08F002-44

CC 38-3 (Plastics Fabrication and Uses)

fuel cell filter press electrolyzer graphite
polymer composite; automobile graphite polymer nanocomposite; free
radical polymn grafting carbon filler graphite composite; bulk
polymn grafting carbon filler graphite composite; polydispersity
index grafting carbon hybrid composite

IT Automobiles

Electrolytic cells

Fuel cells

Hybrid organic-inorganic materials

(method to produce graphite/polymer hybrid composites) IT

2530-85-0, 3-(Trimethoxysilyl) propyl methacrylate 17196-12-2

, 7-Octenyldimethylchlorosilane

RL: RCT (Reactant); RACT (Reactant or reagent)

(free radical graft polymn. in method to produce graphite/polymer hybrid composites)

ANSWER 8 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN L27

ACCESSION NUMBER:

2004:39479 HCAPLUS

DOCUMENT NUMBER:

140:79830

TITLE:

Nonaqueous battery

electrolyte containing a specific Applicant

INVENTOR(S):

Awano, Naomi; Usami, Kyohei; Kubota, Naohiro

PATENT ASSIGNEE(S):

Denso Corporation, Japan; Asahi Denka Co., Ltd.

SOURCE:

U.S. Pat. Appl. Publ., 11 pp.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2004007688	A1	20040115	US 2003-600452	200306
JP 2004039510	A2	20040205	JP 2002-196750	23
EP 1383187	A2	20040121	EP 2003-14518	05
				200307 03

AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK

PRIORITY APPLN. INFO.:

JP 2002-196750

200207

05

OTHER SOURCE(S): MARPAT 140:79830

GI

$$R^{1}$$
 R^{1}
 R^{1}
 R^{3}
 R^{3}

- AB A nonaq. electrolytic soln. comprising an electrolyte salt dissolved in an org. solvent is disclosed. The nonaq. electrolytic soln. contains a silicon compd. represented by formula (I): wherein R1 represents an alkenyl group having 2 to 10 carbon atoms; R2 and R3 each represent an alkyl group having 1 to 10 carbon atoms, an alkoxy group having 1 to 10 carbon atoms, an alkenyl group having 2 to 10 carbon atoms or a halogen atom; and X represents a halogen atom.
- IT 1719-58-0 19304-01-9 38755-76-9 210362-80-4 640269-68-7 640269-69-8 640269-70-1 640269-71-2

RL: MOA (Modifier or additive use); USES (Uses)
 (nonaq. battery electrolyte contg.
 specific silicon compd.)

RN 1719-58-0 HCAPLUS

CN Silane, chloroethenyldimethyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{Cl} \\ | \\ \text{Me-Si-CH----} \text{CH}_2 \\ | \\ \text{Me} \end{array}$$

RN 19304-01-9 HCAPLUS CN Silane, triethenylfluoro- (9CI) (CA INDEX NAME)

$$H_2C = CH - Si - CH = CH_2$$

$$CH = CH_2$$

RN 38755-76-9 HCAPLUS

CN Silane, ethenylfluorodimethyl- (9CI) (CA INDEX NAME)

RN 210362-80-4 HCAPLUS

CN Silane, fluorodimethyl-2-propenyl- (9CI) (CA INDEX NAME)

RN 640269-68-7 HCAPLUS

CN Silane, ethenyldiethylfluoro- (9CI) (CA INDEX NAME)

RN 640269-69-8 HCAPLUS

CN Silane, dibutylethenylfluoro- (9CI) (CA INDEX NAME)

RN 640269-70-1 HCAPLUS

CN Silane, diethenylfluoromethyl- (9CI) (CA INDEX NAME)

MEI HUANG EIC1700 REM4B28 571-272-3952

$$^{\mathrm{Me}}$$
 $^{\mathrm{H}_{2}\mathrm{C}}=\mathrm{CH}-\mathrm{Si}-\mathrm{CH}=\mathrm{CH}_{2}$
 $^{\mathrm{H}_{2}\mathrm{C}}=\mathrm{CH}-\mathrm{Si}-\mathrm{CH}=\mathrm{CH}_{2}$

RN 640269-71-2 HCAPLUS CN Silane, ethenylfluorodimethoxy- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{F} \\ \mid \\ \text{MeO-Si-CH----} \text{CH}_2 \\ \mid \\ \text{OMe} \end{array}$$

IC ICM H01G002-00

INCL 252062200

52-2 (Electrochemical, Radiational, and Thermal Energy Technology)

ST. battery nonag electrolyte specific silicon compd additive

IT Ethers, uses

> RL: DEV (Device component use); USES (Uses) (cyclic; nonaq. battery electrolyte

contg. specific silicon compd.)

ΙT Carboxylic acids, uses

> RL: DEV (Device component use); USES (Uses) (esters; nonaq. battery electrolyte contg. specific silicon compd.)

IT Secondary batteries

> (lithium; nonaq. battery electrolyte contg. specific silicon compd.)

IT Battery electrolytes

(nonaq. battery electrolyte contg.

specific silicon compd.)

IT Amides, uses

Esters, uses

Ethers, uses

Lactones

Sulfones

Sulfoxides

RL: DEV (Device component use); USES (Uses)

(nonaq. battery electrolyte contg.

specific silicon compd.)

IT 96-49-1, Ethylene carbonate 105-58-8, Diethyl carbonate 463-79-6D, Carbonic acid, acyclic compds. 463-79-6D, Carbonic acid, cyclic compds. 616-38-6, Dimethyl carbonate 623-53-0, Ethylmethyl carbonate 4437-85-8, 1,2-Butylene carbonate 7791-03-9, Lithium perchlorate 14283-07-9, Lithium tetrafluoroborate 21324-40-3, Lithium hexafluorophosphate 29935-35-1, Lithium hexafluoroarsenate 33454-82-9, Lithium triflate 90076-65-6 132404-42-3

RL: DEV (Device component use); USES (Uses)

(nonaq. battery electrolyte contq.

specific silicon compd.)

IT 1719-58-0 7440-21-3D, Silicon, compd. 19304-01-9

38755-76-9 210362-80-4 640269-68-7 640269-69-8 640269-70-1 640269-71-2

RL: MOA (Modifier or additive use); USES (Uses)

(nonaq. battery electrolyte contg.

specific silicon compd.)

L27 ANSWER 9 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 200

2003:634130 HCAPLUS

DOCUMENT NUMBER:

139:166967

TITLE:

Anticorrosion coatings for metallic fuel cell components in proton-exchange membrane fuel

cells and direct methanol fuel cells

INVENTOR(S):

Coleman, Ernest A.; Allen, Jeffrey P.

PATENT ASSIGNEE(S):

Gencell Corporation, USA

SOURCE:

PCT Int. Appl., 74 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2003067682	A2	20030814	WO 2003-US3466	
				200302 05
WO 2003067682	A 3	20050616		

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH,

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CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH,
        GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK,
        LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ,
        PL, PT, RO, RU, SD, SE, SG, SK, SL, TJ, TM, TR, TT, TZ, UA,
                        YU, ZA,
        UG, US, UZ, VN,
                                 ZW
    RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,
        BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK,
        EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, SE, SI,
        SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE,
        SN, TD, TG
US 2003170539
                     A1
                            20030911
                                        US 2003-345073
                                                                200301
                                                                15
CA 2474913
                     AA
                            20030814
                                        CA 2003-2474913
                                                                200302
                                                                05
US 2003157391
                     A1
                            20030821
                                        US 2003-358736
                                                                200302
                                                                05
CA 2510358
                     AA
                            20040805
                                        CA 2004-2510358
                                                                200401
                                                                14
WO 2004066470
                     A2
                            20040805
                                        WO 2004-US939
                                                                200401
                                                                14
WO 2004066470
                     A3
                            20041021
        AE, AE, AG, AL, AL, AM, AM, AM, AT, AT, AU, AZ, AZ, BA, BB,
        BG, BG, BR, BR, BW, BY, BY, BZ, BZ, CA, CH, CN, CN, CO, CO,
        CR, CR, CU, CU, CZ, CZ, DE, DE, DK, DK, DM, DZ, EC, EC, EE,
        EE, EG, ES, ES, FI, FI, GB, GD, GE, GE, GH, GM, HR, HR, HU,
        HU, ID, IL, IN, IS, JP, JP, KE, KE, KG, KG, KP, KP, KP, KR,
        KR, KZ, KZ, KZ, LC, LK, LR, LS, LS, LT, LU, LV, MA, MD, MD,
        MG, MK, MN, MW, MX, MX, MZ, MZ, NA, NI
    RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM,
        AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE,
        DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO,
        SE, SI, SK,
                    TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML,
                    TD, TG, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW,
        MR, NE, SN,
        ML, MR, NE, SN,
                        TD, TG
EP 1584119
                            20051012
                     A2
                                        EP 2004-702206
                                                                200401
                                                                14.
        AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,
        PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU,
        SK
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PR'	OR	TTY	APPLN.	INFO.:

US 2002-354554P

200202

05

P

Α

W

US 2003-345073

200301

15

WO 2003-US3466

W

200302

05

WO 2004-US939

200401

14

AB Metallic fuel cell components that are at least partially coated with a coating comprising silane are provided. Methods of protecting a metallic fuel cell component from corrosion is provided, in which the methods comprise at least partially coating a fuel cell bipolar separator plate with a coating comprising a silane. Also included are fuel cells and fuel cell stacks comprising such metallic fuel cell components and methods for manufg. such.

IT 75-94-5, Vinyltrichlorosilane

RL: TEM (Technical or engineered material use); USES (Uses) (anticorrosion coatings for metallic fuel cell components in proton-exchange membrane fuel cells and direct methanol fuel cells)

RN 75-94-5 HCAPLUS

CN Silane, trichloroethenyl- (9CI) (CA INDEX NAME)

 $Cl_3Si-CH=CH_2$

IT 33415-29-1

RL: TEM (Technical or engineered material use); USES (Uses) (panticorrosion coatings for metallic fuel cell components in proton-exchange membrane fuel cells and direct methanol fuel cells)

RN 33415-29-1 HCAPLUS

CN Silane, trichloro-2-propynyl- (8CI, 9CI) (CA INDEX NAME)

$Cl_3Si-CH_2-C \equiv CH$

- IC ICM H01M CC 52-2 (Electrochemical, Radiational, and Thermal Energy Technology) Section cross-reference(s): 38, 55, 56 IT Fuel cells (solid electrolyte, proton-exchange membrane; anticorrosion coatings for metallic fuel cell components in proton-exchange membrane fuel cells and direct methanol fuel cells) 64-19-7, Acetic acid, uses IT 67-63-0, Isopropanol, uses **75-94-5**, Vinyltrichlorosilane 78-08-0, 78-10-4, Tetraethoxysilane Vinyltriethoxysilane 108-88-3, Toluene, uses 681-84-5, Tetramethoxysilane 919-30-2, 3-Aminopropyltriethoxysilane 999-97-3, Hexamethyldisilazane 1185-55-3, Methyltrimethoxysilane 1330-20-7, Xylene, 1067-53-4 1760-24-3, N-(2-Aminoethyl)-3-aminopropyltrimethoxysilane 2530-83-8, 3-Glycidoxypropyltrimethoxysilane 2530-85-0, γ-Methacryloxypropyltrimethoxysilane 2768-02-7, Vinyltrimethoxysilane 3069-42-9, Octadecyltrimethoxysilane 3388-04-3, 2-(3,4-Epoxycyclohexyl)-ethyltrimethoxysilane 4130-08-9, Vinyl triacetoxysilane 4420-74-0, 3-Mercaptopropyltrimethoxysilane 4766-57-8, Tetra-n-butoxysilane 5700-28-7 7538-45-6, 2-Mercaptoethyltrimethoxysilane 7803-62-5D. Silane, 1-cyanobutyltrialkoxy deriv. 7803-62-5D, Silane, 1-cyanoisobutyltrialkoxy deriv. 7803-62-5D, Silane, cyanoethyltrialkoxy deriv. 7803-62-5D, Silane, cyanoisobutyltrialkoxy deriv. 7803-62-5D, Silane, cyanophenyltrialkoxy deriv. cyanopropyltrialkoxy deriv. 7803-62-5D, Silane, 13822-56-5, 3-Aminopropyltrimethoxysilane 14044-97-4, Mercaptosilane
 - γ-Ureidopropyltriethoxysilane 25904-66-9 32957-40-7, Ethynyl trimethoxysilane 51473-59-7 575464-53-8 57

Ethynyl trimethoxysilane 51473-59-7 575464-53-8 575464-54-9 RL: TEM (Technical or engineered material use); USES (Uses)

17053-34-8

23779-32-0.

20208-39-3,

(anticorrosion coatings for metallic fuel cell components in proton-exchange membrane fuel cells and direct methanol fuel cells)

IT 33415-29-1

RL: TEM (Technical or engineered material use); USES (Uses) (panticorrosion coatings for metallic fuel cell components in proton-exchange membrane fuel cells and direct methanol fuel cells)

16881-77-9, Methyldimethoxysilane

γ-Acryloxypropyltriethoxysilane

L27 ANSWER 10 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2002:499574 HCAPLUS

DOCUMENT NUMBER:

137:35458

TITLE:

Crosslinked or modified polymeric porous films

as separators for batteries with

nonaqueous electrolytes

INVENTOR(S):

Kenichiro, Kami; Hiroshi, Ueshima; Ryuichirou, Shinkai; Norikazu, Hosokawa; Manabu, Yamada;

Hideo, Amaki; Tomoaki, Tamura

PATENT ASSIGNEE(S):

Denso Corp., Japan

SOURCE:

Fr. Demande, 75 pp. CODEN: FRXXBL

DOCUMENT TYPE:

Patent

LANGUAGE:

French

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
FR 2814284	A1	20020322	FR 2001-12110	
				200109 19
JP 2002170541	A2	20020614	JP 2001-145341	10
				200105 15
PRIORITY APPLN. INFO.:			JP 2000-287145 A	
				200009
				21
			JP 2001-145341 A	
				200105 15
			JP 2000-141476 A	200005
				200005 15

AB Non-swelling polymeric separators for batteries with non-aq. electrolytes are prepd. by crosslinking or modifying a porous polymer film with compds. that formed the crosslinks or added chains by at least 2 carbon atoms away from the linking units between the monomers (e.g., the ester linkage of polyesters). The polymeric film can be selected from

polybenzimidazoles, polyimides, polyether-polyimides, polyamide-polyimides, polyphenylene sulfides, polyphenylene oxides, polyether-polysulfones, polysulfones, polyether-polyketones, aramides, satd. polyesters, polyoxymethylenes, etc. Suitable crosslinking agents or modifiers include acrylates, vinyl compds., and functionalized (unsatd.) alkoxysilanes.

IT 75-94-5, Vinyltrichlorosilane

RL: DEV (Device component use); RCT (Reactant); RACT (Reactant or reagent); USES (Uses)

(crosslinking or modifying agent; polymeric porous films as separators for batteries with nonaq.

electrolytes)

RN 75-94-5 HCAPLUS

CN Silane, trichloroethenyl- (9CI) (CA INDEX NAME)

Cl₃Si-CH=CH₂

- IC H01M010-38
- CC **52-2** (Electrochemical, Radiational, and Thermal Energy Technology)
 Section cross-reference(s): 38
- ST nonswelling battery separator nonaq
 electrolyte; crosslinking modification polymeric
 battery separator nonaq electrolyte;
 polyester crosslinked film battery separator nonaq
 electrolyte
- IT Polysiloxanes, uses

RL: DEV (Device component use); RCT (Reactant); RACT (Reactant or reagent); USES (Uses)

(acrylic, crosslinking or modifying agent; polymeric porous films as separators for batteries with nonaq.

electrolytes)

IT Polyamide fibers, uses

RL: DEV (Device component use); RCT (Reactant); RACT (Reactant or reagent); USES (Uses)

(aramid, crosslinked or modified, battery separators; polymeric porous films as separators for batteries with nonag. electrolytes)

IT Polyesters, uses

RL: DEV (Device component use); RCT (Reactant); RACT (Reactant or reagent); USES (Uses)

(arom., Vylon KS001, crosslinked or modified, battery separators; polymeric porous films as separators for

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batteries with nonaq. electrolytes)
IT
     Primary battery separators
     Secondary battery separators
        (crosslinked or modified polymeric porous films as separators for
        batteries with nonag. electrolytes)
IT
     Fluoropolymers, uses
     Polyamides, uses
     Polybenzimidazoles
     Polyesters, uses
     Polyimides, uses
     Polyoxymethylenes, uses
     Polyoxyphenylenes
     Polysulfones, uses
     Polythiophenylenes
     RL: DEV (Device component use); RCT (Reactant); RACT (Reactant or
     reagent); USES (Uses)
        (crosslinked or modified, battery separators; polymeric
        porous films as separators for batteries with
        nonaq. electrolytes)
IT
     Battery electrolytes
        (nonaq.; crosslinked or modified polymeric porous films
        as separators for batteries with nonag.
        electrolytes)
IT
     Polyimides, uses
     RL: DEV (Device component use); RCT (Reactant); RACT (Reactant or
     reagent); USES (Uses)
        (polyamide-, crosslinked or modified, battery
        separators; polymeric porous films as separators for
        batteries with nonag. electrolytes)
IT
     Polyimides, uses
     Polyketones
     Polysulfones, uses
     RL: DEV (Device component use); RCT (Reactant); RACT (Reactant or
     reagent); USES (Uses)
        (polyether-, crosslinked or modified, battery
        separators; polymeric porous films as separators for
        batteries with nonaq. electrolytes)
IT
     Polyamides, uses
     Polyethers, uses
     RL: DEV (Device component use); RCT (Reactant); RACT (Reactant or
     reagent); USES (Uses)
        (polyimide-, crosslinked or modified, battery
        separators; polymeric porous films as separators for
        batteries with nonag. electrolytes)
IT
     Polyethers, uses
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RL: DEV (Device component use); RCT (Reactant); RACT (Reactant or reagent); USES (Uses)

(polyketone-, crosslinked or modified, battery

separators; polymeric porous films as separators for

batteries with nonag. electrolytes)

IT Acrylic polymers, uses

RL: DEV (Device component use); RCT (Reactant); RACT (Reactant or reagent); USES (Uses)

(polysiloxane-, crosslinking or modifying agent; polymeric porous films as separators for batteries with nonag.

electrolytes)

IT Polyethers, uses

RL: DEV (Device component use); RCT (Reactant); RACT (Reactant or reagent); USES (Uses)

(polysulfone-, crosslinked or modified, **battery** separators; polymeric porous films as separators for

batteries with nonaq. electrolytes)

IT 2530-85-0, γ-(Methacryloxypropyl)trimethoxysilane

RL: DEV (Device component use); RCT (Reactant); RACT (Reactant or reagent); USES (Uses)

(KBM503, crosslinking or modifying agent; polymeric porous films as separators for **batteries** with **nonaq**.

electrolytes)

IT 4369-14-6, 2-Propenoic acid, 3-(trimethoxysilyl)propyl ester RL: DEV (Device component use); RCT (Reactant); RACT (Reactant or reagent); USES (Uses)

(KBM5103, crosslinking or modifying agent; polymeric porous films as separators for batteries with nonaq.

electrolytes)

IT 1025-15-6, Triallyl isocyanurate

RL: DEV (Device component use); RCT (Reactant); RACT (Reactant or reagent); USES (Uses)

(TAIC, crosslinking or modifying agent; polymeric porous films as separators for **batteries** with **nonaq**.

electrolytes)

9016-80-2, Poly(methylpentene) 9020-73-9, Polyethylene naphthalate 24937-79-9, Poly(vinylidene fluoride) 24968-11-4, Polyethylene naphthalate 24968-12-5, Polybutylene terephthalate 25038-59-9, Polyethylene terephthalate, uses 26062-94-2, Polybutylene terephthalate 28779-82-0, Polybutylene naphthalate 51806-50-9, Polybutylene naphthalate

RL: DEV (Device component use); RCT (Reactant); RACT (Reactant or reagent); USES (Uses)

(crosslinked or modified, battery separators; polymeric porous films as separators for batteries with

nonaq. electrolytes)

IT

75-94-5, Vinyltrichlorosilane 78-08-0, Vinyltriethoxysilane 97-90-5, Ethylene glycol dimethacrylate 131-17-9, Diallyl phthalate 1067-53-4, Vinyltris(βmethoxyethoxy) silane 1321-74-0, Divinylbenzene, uses 1337-81-1, Vinylpyridine 2768-02-7, Vinyltrimethoxysilane 3030-60-2, Allyl 3290-92-4, Trimethylolpropane trimethacrylate isocyanurate 6294-79-7, Diallyl isocyanurate 21142-29-0, γ-(Methacryloxypropyl) triethoxysilane 25013-15-4, Vinyltoluene

65100-04-1, γ-(Methacryloxypropyl)methyldiethoxysilane RL: DEV (Device component use); RCT (Reactant); RACT (Reactant or reagent); USES (Uses)

(crosslinking or modifying agent; polymeric porous films as separators for batteries with nonag. electrolytes)

ANSWER 11 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:598346 HCAPLUS

DOCUMENT NUMBER:

135:155269

TITLE: Nonaqueous electrolyte:

batteries and their manufacture

INVENTOR(S): Yoshida, Yasuhiro; Hiroi, Osamu; Nakao,

Yukiyasu; Shiota, Hisashi; Aihara, Shigeru; Takemura, Daigo; Urushibata, Hiroaki; Murai, Michio; Nishimura, Takashi; Aragane, Jun;

Kurata, Tetsuyuki

PATENT ASSIGNEE(S): Mitsubishi Denki K. K., Japan

SOURCE: PCT Int. Appl., 23 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001059871	A1	20010816	WO 2000-JP736	200002
W: JP, KR, US RW: DE, FR EP 1184927	A1	20020306	EP 2000-902897	200002

R: DE, FR PRIORITY APPLN. INFO.:

WO 2000-JP736

W

200002 10

AB The **batteries** have their electrodes and/or separators coated with a mixt. contg. **nonaq.** solvents, fine inorg. particles, and ≥1 of dewatering agents, hydrophobic agents, silylation agents, and coupling agents; and are prepd. by applying the mixt. on the electrodes and/or separators.

TT 75-94-5, Vinyl trichlorosilane
RL: NUU (Other use, unclassified); USES (Uses)
(coupling agents in manuf. of secondary lithium batteries
with inorg. powder coated electrodes and separators)

RN 75-94-5 HCAPLUS

CN Silane, trichloroethenyl- (9CI) (CA INDEX NAME)

Cl₃Si-CH=CH₂

IC ICM H01M010-40 ICS H01M004-04; H01M006-14; H01M002-14

CC **52-2** (Electrochemical, Radiational, and Thermal Energy Technology)

ST nonaq battery electrode separator hydrophobic coating; inorg powder coating nonaq battery electrode separator

IT Secondary batteries

(lithium; manuf. of secondary lithium batteries with inorg. powder coated electrodes and separators)

IT Battery electrodes

Secondary battery separators

(manuf. of secondary lithium batteries with inorg. powder coated electrodes and separators)

TT 75-94-5, Vinyl trichlorosilane 999-97-3,
Hexamethyldisilazane 2530-85-0, γ(Methacryloxypropyl)trimethoxysilane 3068-76-6,
N-Phenyl-γ-aminopropyltrimethoxysilane
RL: NUU (Other use, unclassified); USES (Uses)

(coupling agents in manuf. of secondary lithium batteries with inorg. powder coated electrodes and separators)

IT 7440-44-0, Carbon, uses 9003-07-0, Polypropylene 12190-79-3,
 Cobalt lithium oxide (CoLiO2)
 RL: DEV (Device component use); PEP (Physical, engineering or

chemical process); PROC (Process); USES (Uses)
 (manuf. of secondary lithium batteries with inorg.
 powder coated electrodes and separators)

IT 1344-28-1, Alumina, uses

RL: MOA (Modifier or additive use); USES (Uses) (manuf. of secondary lithium batteries with inorg. powder coated electrodes and separators)

IT 68-12-2, DMF, uses

RL: NUU (Other use, unclassified); USES (Uses)
(nonaq. solvent in manuf. of secondary lithium
batteries with inorg. powder coated electrodes and
separators)

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L27 ANSWER 12 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2001:270894 HCAPLUS

DOCUMENT NUMBER:

135:61379

TITLE:

Electrochemical Synthesis of Functional Aryland Heteroarylchlorosilanes. Application to the Preparation of Donor-Acceptor or Donor-Donor

Organosilicon Molecules

AUTHOR (S):

Moreau, Carole; Serein-Spirau, Francoise;

Bordeau, Michel; Biran, Claude

CORPORATE SOURCE:

Laboratoire de Chimie Organique et

Organometallique, UMR 5802 CNRS Universite

Bordeaux 1, Talence, F-33405, Fr.

SOURCE:

Organometallics (2001), 20(10), 1910-1917

CODEN: ORGND7; ISSN: 0276-7333

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 135:61379

AB 44A variety of functional aryl- and heteroarylchlorosilanes were prepd. by electrochem. redn. of halothiophenes, halofurans, halopyridines, and substituted aryl halides in the presence of a large excess of com. organodi- or trichlorosilanes using an undivided cell, a sacrificial Mg or Al anode, a const. c.d., and tetrabutylammonium bromide as the supporting electrolyte.

New structures are described, and some examples of the use of these more elaborate chlorosilanes are given, particularly as interesting precursors for the prepn. of polarized D-A and D-D organosilicon models. Thus, electrolysis of 2-bromothiophene contg. Mg or Al anode in the presence of THF/HMPA/Bu4NBr and Me2SiCl2 gave 88%

2-thienylchlorodimethylsilane. IT 75-94-5, Vinyltrichlorosilane 124-70-9, Dichloro (methyl) vinylsilane RL: RCT (Reactant); RACT (Reactant or reagent) (electrochem. synthesis of functional aryl- and heteroarylchlorosilanes. and application to prepn. of donor-acceptor or donor-donor organosilicon mols.) RN 75-94-5 HCAPLUS Silane, trichloroethenyl- (9CI) (CA INDEX NAME) CN $Cl_3Si-CH=CH_2$ RN 124-70-9 HCAPLUS CN Silane, dichloroethenylmethyl- (9CI) (CA INDEX NAME) Cl Me-Si-CH=CH2 Cl 29-6 (Organometallic and Organometalloidal Compounds) Section cross-reference(s): 21, 72 75-78-5, Dichlorodimethylsilane 75-79-6, Methyltrichlorosilane IT 75-94-5, Vinyltrichlorosilane 96-43-5, 2-Chlorothiophene 100-00-5, 1-Chloro-4-nitrobenzene 109-04-6, 2-Bromopyridine **124-70-9**, Dichloro (methyl) vinylsilane 149-74-6, Dichloro (methyl) phenylsilane 402-43-7, 1-Bromo-4trifluoromethylbenzene 460-00-4, 1-Bromo-4-fluorobenzene 584-12-3, 2-Bromofuran 586-77-6, 1-Bromo-4-dimethylaminobenzene 586-78-7, 1-Bromo-4-nitrobenzene 626-55-1, 3-Bromopyridine 1003-09-4, 2-Bromothiophene 3141-27-3, 2,5-Dibromothiophene RL: RCT (Reactant); RACT (Reactant or reagent) (electrochem. synthesis of functional aryl- and heteroarylchlorosilanes. and application to prepn. of donor-acceptor or donor-donor organosilicon mols.) REFERENCE COUNT: 38 THERE ARE 38 CITED REFERENCES AVAILABLE

IN THE RE FORMAT

L27 ANSWER 13 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

FOR THIS RECORD. ALL CITATIONS AVAILABLE

ACCESSION NUMBER: 2000:131373 HCAPLUS

DOCUMENT NUMBER: 132:200258

TITLE: Effect of surface modifications on spectral

shift of electroluminescence of porous n-Si in

S2082- under cathodic bias

AUTHOR(S): Lee, Hyun-Gon; Park, Heung-Shik; Lim, Hyun-Eui;

Lee, Yeonhee; Kim, Kang-Jin

CORPORATE SOURCE: Department of Chemistry, Korea University,

Seoul, 136-701, S. Korea

SOURCE: Journal of the Electrochemical Society (2000),

147(2), 650-654

CODEN: JESOAN; ISSN: 0013-4651

PUBLISHER: Electrochemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

The spectral shifts of visible electroluminescence from chem. modified porous Si under cathodic bias in an electrolyte contg. S2082- were studied by luminescence and FTIR measurements. Modification of porous Si with sol. polyaniline and silane compds. results in blueshifts, whereas red shifts are obsd. with H+ and O+ ion implantations. Based on a recently proposed mechanism that silicon hydride acts as an electron injector to sulfate radical anion, the blueshift is linked to the lowering of the energy levels of silicon hydrides on the porous Si surface with the bandedges being invariant with the chem. modifications. The red shift is seen as a result of the conversion of silicon dihydride to silicon monohydride, with the energy levels of silicon monohydride lying higher than silicon dihydride.

IT 1719-58-0, Chlorodimethylvinylsilane

RL: DEV (Device component use); PEP (Physical, engineering or chemical process); PRP (Properties); PROC (Process); USES (Uses) (spectral shifts of visible electroluminescence from vapor phase chlorodimethylvinylsilane modified porous Si under cathodic bias in electrolyte contq. S2082-)

RN 1719-58-0 HCAPLUS

CN Silane, chloroethenyldimethyl- (9CI) (CA INDEX NAME)

- CC 72-2 (Electrochemistry)
 Section cross-reference(s): 36, 73
- IT 12408-02-5, Hydrogen ion, uses
 RL: MOA (Modifier or additive use); PRP (Properties); USES (Uses)
 (electroluminescence of H+-implanted porous Si under cathodic bias in electrolyte contg. S2082-)
- IT 14581-93-2, Oxygen, ion O1+), uses
 RL: MOA (Modifier or additive use); PRP (Properties); USES (Uses)
 (electroluminescence of O1+-implanted porous Si under cathodic bias in electrolyte contq. S2O82-)
- IT 18162-84-0, Chlorodimethyloctylsilane
 RL: DEV (Device component use); PEP (Physical, engineering or chemical process); PRP (Properties); PROC (Process); USES (Uses)
 (spectral shifts of visible electroluminescence from vapor phase chlorodimethyloctylsilane modified porous Si under cathodic bias in electrolyte contg. S2082-)
- IT 1719-58-0, Chlorodimethylvinylsilane
 RL: DEV (Device component use); PEP (Physical, engineering or
 chemical process); PRP (Properties); PROC (Process); USES (Uses)
 (spectral shifts of visible electroluminescence from vapor phase
 chlorodimethylvinylsilane modified porous Si under cathodic bias
 in electrolyte contq. S2082-)
- IT 18419-53-9, Chlorodiphenylvinylsilane
 RL: DEV (Device component use); PEP (Physical, engineering or chemical process); PRP (Properties); PROC (Process); USES (Uses)
 (spectral shifts of visible electroluminescence from vapor phase chlorodiphenylvinylsilane modified porous Si under cathodic bias in electrolyte contg. S2082-)
- TT 75-77-4, Chlorotrimethylsilane, uses
 RL: DEV (Device component use); PEP (Physical, engineering or chemical process); PRP (Properties); PROC (Process); USES (Uses)
 (spectral shifts of visible electroluminescence from vapor phase chlorotrimethylsilane modified porous Si under cathodic bias in

electrolyte contg. S2082-)

REFERENCE COUNT:

38 THERE ARE 38 CITED REFERENCES AVAILABLE

FOR THIS RECORD. ALL CITATIONS AVAILABLE

IN THE RE FORMAT

L27 ANSWER 14 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1999:390273 HCAPLUS

DOCUMENT NUMBER:

131:21352

TITLE:

Manufacture of electrolytes and

secondary batteries

INVENTOR(S):

Yamamoto, Tomoya; Kawakami, Soichiro

PATENT ASSIGNEE(S):

Canon K. K., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 14 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 11162508	A2	19990618	JP 1998-271586	199809 25
JP 3428910	B2	20030722		
US 6277525	B1	20010821	US 1998-159572 .	199809 24
PRIORITY APPLN. INFO.:			JP 1997-259996 A	199709 25

AB The org. F and Si contg. salt electrolytes are prepd. by reacting at least an org. silane with a fluoro compd. in a nonag. solvent based medium. Secondary Li batteries are prepd. by using the electrolytes.

IT 226724-11-4P

RL: IMF (Industrial manufacture); PEP (Physical, engineering or chemical process); PREP (Preparation); PROC (Process) (intermediates in manuf. of org. fluorine and silicon contg.

salts for electrolytes for secondary lithium

batteries)

RN 226724-11-4 HCAPLUS

CN Ethanaminium, N,N,N-triethyl-, (OC-6-21)-

ethenylpentafluorosilicate(2-) (2:1) (9CI) (CA INDEX NAME)

CM 1

CRN 27900-02-3 CMF C2 H3 F5 Si

CCI CCS

CM 2

CRN 66-40-0 CMF C8 H20 N

IT 226724-03-4P

RL: DEV (Device component use); IMF (Industrial manufacture); PREP (Preparation); USES (Uses)

(manuf. of org. fluorine and silicon contg. salts for electrolytes for secondary lithium batteries)

RN 226724-03-4 HCAPLUS

CN Silicate(2-), ethenylpentafluoro-, dilithium, (OC-6-21)- (9CI) (CA INDEX NAME)

●2 Li+

IC ICM H01M010-40

CC **52-2** (Electrochemical, Radiational, and Thermal Energy Technology)

ST secondary lithium battery electrolyte manuf; fluoro silicon salt electrolyte lithium battery

IT Battery electrolytes

(manuf. of org. fluorine and silicon contg. salts for electrolytes for secondary lithium batteries)

TT 76-86-8, Triphenyl chlorosilane 78-08-0, Vinyl triethoxysilane 80-10-4 429-41-4, Tetrabutylammonium fluoride 665-46-3, Tetraethylammonium fluoride 780-69-8, Phenyl triethoxy silane 3027-21-2, Dimethoxy methyl phenyl silane 7789-24-4, Lithium fluoride, reactions 12125-01-8, Ammonium fluoride ((NH4)F) 17689-77-9

RL: RCT (Reactant); RACT (Reactant or reagent)
(in manuf. of org. fluorine and silicon contg. salts for
electrolytes for secondary lithium batteries)

IT 91811-92-6P 189008-37-5P 226724-05-6P 226724-06-7P 226724-07-8P **226724-11-4P**

RL: IMF (Industrial manufacture); PEP (Physical, engineering or chemical process); PREP (Preparation); PROC (Process) (intermediates in manuf. of org. fluorine and silicon contg. salts for electrolytes for secondary lithium

electrolytes for secondary lithium batteries)

60-29-7, Diethyl ether, uses 67-68-5, Dmso, uses IT75-05-8, Acetonitrile, uses 78-93-3, Ethyl methyl ketone, uses 108-90-7, Chlorobenzene, uses

RL: NUU (Other use, unclassified); USES (Uses)

(solvents in manuf. of org. fluorine and silicon contg. salts for electrolytes for secondary lithium batteries)

ANSWER 15 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1998:715934 HCAPLUS

DOCUMENT NUMBER:

129:333323

TITLE:

Carbonaceous active material for lithium-ion

battery anodes and its manufacture

INVENTOR(S):

Barker, Jeremy; Koksbang, Rene

PATENT ASSIGNEE(S):

Valence Technology Inc, USA

SOURCE:

U.S., 10 pp. CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5830602	A	19981103	US 1997-802977	
PRIORITY APPLN. INFO.:			US 1997-802977	199702 20
TRIORITI AFFIIN. INFO			05 1997-002977	199702 20

- AB The active material contg. a dopant is prepd. by providing vinyl monomers CH2CH(G), where G represents a pendant group comprising ≥1 element different from C; electropolymg. the monomers; and heating the obtained polymer to pyrolyze the polymer. In the obtained carbonaceous active material, C constitutes the largest wt. portion and the ≥1 element constitutes the 2nd largest wt. portion.
- IT 75-94-5, Vinyltri (chloro) silane

RL: NUU (Other use, unclassified); USES (Uses)

(in manuf. of carbonaceous active material for lithium-ion battery anodes)

75-94-5 HCAPLUS RN

Silane, trichloroethenyl- (9CI) (CA INDEX NAME) CN

 $Cl_3Si-CH=CH_2$

IC ICM H01M004-58

INCL 429218000

CC 52-2 (Electrochemical, Radiational, and Thermal Energy Technology) Section cross-reference(s): 38, 72

ST carbonaceous active material lithium ion battery; battery anode lithium ion carbonaceous material

IT Battery anodes

(dopant-contg. carbonaceous active material for lithium-ion)

IT Carbonaceous materials (technological products)

RL: DEV (Device component use); SPN (Synthetic preparation); PREP (Preparation); USES (Uses)

(lithium-ion battery anodes from dopant-contg.)

TT 75-94-5, Vinyltri(chloro)silane 77-77-0, Vinyl sulfone 88-12-0, 1-Vinyl-2-pyrrolidinone, uses 100-43-6, 4-Vinylpyridine 100-69-6, 2-Vinylpyridine 105-38-4, Vinyl propionate 754-05-2, Vinyltri(methyl)silane 2768-02-7, Vinyltri(methoxy)silane 3485-84-5, n-Vinylphthalimide 4177-16-6, 2-Vinylpyrazine 18666-68-7, Vinyltri(phenyl)silane 29383-23-1, Vinylimidazole 56512-51-7, Vinylqlycine

RL: NUU (Other use, unclassified); USES (Uses)
(in manuf. of carbonaceous active material for lithium-ion battery anodes)

REFERENCE COUNT:

THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L27 ANSWER 16 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

25

ACCESSION NUMBER:

1998:594806 HCAPLUS

DOCUMENT NUMBER:

129:247668

TITLE:

Hydrogen-absorbing alloy powders surface treated

with coupling agents and manufacture of

hydrogen-absorbing anodes

INVENTOR(S):

Ikishima, Kenji; Kaminaka, Hideya

PATENT ASSIGNEE(S):

Sumitomo Metal Industries, Ltd., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 7 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent Japanese

LANGUAGE:

r. 1

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
 JP 10241678	A2	19980911	`JP 1997-43891	
·				199702 27
PRIORITY APPLN. INFO.:			JP 1997-43891	
				199702 27

AB Claimed H-absorbing alloy powders are surface treated with silane-based or Ti-based coupling agents. The title anodes are manufd. by coating substrates with slurries or pastes contg. binders and dispersed with the above alloy powders and then rolling the coated materials. Resulting batteries have high filling d. and discharge capacity and prevent exfoliation of active mass.

IT 75-94-5, Vinyltrichlorosilane

RL: MOA (Modifier or additive use); USES (Uses)
(addn. of silane- or titanium-type coupling agents in manuf. of hydrogen-absorbing alloy anodes for batteries)

RN 75-94-5 HCAPLUS

CN Silane, trichloroethenyl- (9CI) (CA INDEX NAME)

$Cl_3Si-CH=CH_2$

- IC ICM H01M004-38
 - ICS B22F001-00; C22C019-00; H01M004-26; H01M004-62
- CC 52-2 (Electrochemical, Radiational, and Thermal Energy Technology) Section cross-reference(s): 56
- ST coupling agent hydrogen absorbing anode **battery**; silane coupling agent hydrogen absorbing alloy; titanium coupling agent hydrogen absorbing alloy
- IT Battery anodes
 - Coupling agents

(addn. of silane- or titanium-type coupling agents in manuf. of hydrogen-absorbing alloy anodes for **batteries**)

IT 181147-99-9

RL: DEV (Device component use); PEP (Physical, engineering or chemical process); TEM (Technical or engineered material use); PROC (Process); USES (Uses)

(addn. of silane- or titanium-type coupling agents in manuf. of hydrogen-absorbing alloy anodes for **batteries**)

T75-94-5, Vinyltrichlorosilane 78-08-0,
Vinyltriethoxysilane 919-30-2, γ-Aminopropyltriethoxysilane
1760-24-3, N-(β-Aminoethyl)-γ-aminopropyltrimethoxysilane
2530-87-2, γ-Chloropropyltrimethoxysilane 7787-93-1
18171-19-2, γ-Chloropropylmethyldimethoxysilane 61417-49-0,
Isopropyltriisostearoyltitanate 61436-47-3, Isopropyl
tris(dioctylphosphato)titanate 64060-97-5 64157-14-8,
Tetra(2,2-diallyloxymethyl-1-butyl)bis(di-tridecyl)phosphitetitanate
90959-84-5 91858-93-4
RL: MOA (Modifier or additive use); USES (Uses)

(addn. of silane- or titanium-type coupling agents in manuf. of hydrogen-absorbing alloy anodes for **batteries**)

IT 1333-74-0, Hydrogen, uses

RL: DEV (Device component use); USES (Uses)
(alloys contg. absorbed; addn. of silane- or titanium-type
coupling agents in manuf. of hydrogen-absorbing alloy anodes for
batteries)

L27 ANSWER 17 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1998:358251 HCAPLUS

DOCUMENT NUMBER: 129:97638

TITLE: ORMOCERs as inorganic-organic

The state of the s

electrolytes for new solid state lithium

batteries and supercapacitors

AUTHOR(S): Popall, M.; Andrei, M.; Kappel, J.; Kron, J.;

Olma, K.; Olsowski, B.

CORPORATE SOURCE: Fraunhofer-Inst. Silicatforschung, Wurzburg,

D-97082, Germany

SOURCE: Electrochimica Acta (1998), 43(10-11), 1155-1161

CODEN: ELCAAV; ISSN: 0013-4686

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

ORMOCERS (ORganically Modified CERamics) are inorg.-org. copolymers which are synthesized as matrix for Li-ion conduction. The inorg. oxidic backbone of these materials results from polycondensation of alkoxy compds. whereas the org. network is formed from reactive functional groups R' of alkoxysilanes of the type R'Si(OR)3, or by co-polymg. reactive org. monomers with reactive functionalized alkoxysilanes. Depending on the reactive org. functionalities and their thermal and UV-initiated org. crosslinking reactions the materials were adapted to the needs of battery and supercapacitor manufg. For ionic cond. polyethers with different chain lengths and functionalized (e.g. epoxy) termination sites were synthesized and attached to organically functionalized oxidic

oligomers. Conductivities of up to 10-4 Ω -1 cm-1 at room temp. were achieved without plasticizer. The **electrolytes** form an amorphous network with configuration temps. (according to Vogel-Tammann-Fulcher) close to -80°, several degrees below the transformation temp. (measured by DSC) in agreement with conventional configuration theory. The activation energies correlate favorably with results for good polymer **electrolytes.**

IT 1871-21-2, Chlorotrivinylsilane

RL: RCT (Reactant); RACT (Reactant or reagent)
(organically modified ceramics as inorg.-org.
electrolytes for new solid state lithium
batteries and supercapacitors)

RN 1871-21-2 HCAPLUS

CN Silane, chlorotriethenyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{Cl} \\ | \\ \text{H}_2\text{C} = \text{CH} - \text{Si} - \text{CH} = \text{CH}_2 \\ | \\ \text{CH} = \text{CH}_2 \end{array}$$

CC 52-2 (Electrochemical, Radiational, and Thermal Energy Technology) Section cross-reference(s): 38, 57, 76

ST battery supercapacitor electrolyte organically modified ceramic

IT Polyoxyalkylenes, preparation

RL: SPN (Synthetic preparation); PREP (Preparation) (electrolyte contg.; organically modified ceramics as inorg.-org. electrolytes for new solid state lithium batteries and supercapacitors)

IT Polysiloxanes, preparation

Polysiloxanes, preparation

RL: SPN (Synthetic preparation); PREP (Preparation) (epoxy; organically modified ceramics as inorg.-org. electrolytes for new solid state lithium batteries and supercapacitors)

IT Secondary batteries

(lithium; organically modified ceramics as inorg.-org. electrolytes for new solid state lithium

batteries and supercapacitors)

IT Battery electrolytes
Ceramics

Electric conductivity Hydrolysis Ionic conductivity (organically modified ceramics as inorg.-org. electrolytes for new solid state lithium batteries and supercapacitors) IT Epoxy resins, preparation Epoxy resins, preparation RL: SPN (Synthetic preparation); PREP (Preparation) (polysiloxane-; organically modified ceramics as inorg.-org. electrolytes for new solid state lithium batteries and supercapacitors) IT Capacitors (super-; organically modified ceramics as inorq.-org. electrolytes for new solid state lithium batteries and supercapacitors) IT 7791-03-9P, Lithium perchlorate 25322-68-3P, Peo RL: SPN (Synthetic preparation); PREP (Preparation) (electrolyte contg.; organically modified ceramics as inorg.-org. electrolytes for new solid state lithium batteries and supercapacitors) IT 12125-01-8, Ammonium fluoride RL: CAT (Catalyst use); USES (Uses) (organically modified ceramics as inorg.-org. electrolytes for new solid state lithium batteries and supercapacitors) IT 1871-21-2, Chlorotrivinylsilane 2530-83-8, 3-Glycidyloxypropyltrimethoxysilane RL: RCT (Reactant); RACT (Reactant or reagent) (organically modified ceramics as inorg.-org. electrolytes for new solid state lithium batteries and supercapacitors) IT 56325-93-0P, 3-Glycidyloxypropyltrimethoxysilane homopolymer RL: SPN (Synthetic preparation); PREP (Preparation) (organically modified ceramics as inorg.-org. electrolytes for new solid state lithium batteries and supercapacitors) REFERENCE COUNT: THERE ARE 17 CITED REFERENCES AVAILABLE 17 FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L27 ANSWER 18 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

1998:358225 HCAPLUS

Towards solid state lithium batteries

129:97635

ACCESSION NUMBER:

DOCUMENT NUMBER:

TITLE:

based on ORMOCER electrolytes

AUTHOR(S): Skaarup, Steen; West, Keld; Zachau-Christiansen,

Birgit; Popall, Michael; Kappel, Jurgen; Kron,

Johanna; Eichinger, Gunther; Semrau, Gunther

CORPORATE SOURCE: Departments of Physics and of Chemistry,

Technical University of Denmark, Lyngby,

DK-2800, Den.

SOURCE: Electrochimica Acta (1998), 43(10-11), 1589-1592

CODEN: ELCAAV; ISSN: 0013-4686

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

AB ORMOCER polymer electrolytes have been tested both as separator electrolytes, and as binder electrolyte in composite cathodes of lithium secondary batteries. The interface stability towards metallic lithium has been strongly improved by careful control of purity and conditions during polymn. About 900 cycles have been obtained with utilizations decreasing from 65 to 25% with the total discharge capacity corresponding to 330 full discharges. The charge factor during long term cycling is very close to 1 (1.004 ± 0.006), indicating that the amt. of parasitic side reactions can be minimal.

IT 1871-21-2, ChlorotriVinylsilane 4028-23-3, Allyl

dimethylchlorosilane

RL: RCT (Reactant); RACT (Reactant or reagent)
 (solid state lithium batteries based on organically
 modified ceramic electrolytes)

RN 1871-21-2 HCAPLUS

CN Silane, chlorotriethenyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{Cl} \\ | \\ \text{H}_2\text{C} = \text{CH} - \text{Si} - \text{CH} = \text{CH}_2 \\ | \\ \text{CH} = \text{CH}_2 \end{array}$$

RN 4028-23-3 HCAPLUS

CN Silane, chlorodimethyl-2-propenyl- (9CI) (CA INDEX NAME)

```
C1
Me-Si-CH_2-CH=CH_2
   Me
CC
     52-2 (Electrochemical, Radiational, and Thermal Energy Technology)
     Section cross-reference(s): 38, 57
ST
     battery electrolyte separator organically
     modified ceramic
     Secondary batteries
IT
        (lithium; solid state lithium batteries based on
        organically modified ceramic electrolytes)
IT
     Battery electrolytes
     Ceramic composites
     Electric impedance
     Hydrolysis
     Secondary battery separators
     Sol-gel processing
        (solid state lithium batteries based on organically
        modified ceramic electrolytes)
IT
     Polyoxyalkylenes, uses
     RL: MOA (Modifier or additive use); USES (Uses)
        (solid state lithium batteries based on organically
        modified ceramic electrolytes)
IT
     12162-79-7, Lithium manganese oxide limno2
     RL: DEV (Device component use); USES (Uses)
        (cathodes; solid state lithium batteries based on
        organically modified ceramic electrolytes)
IT
     24991-55-7, Polyethyleneglycol dimethyl ether
                                                      25322-68-3
     RL: MOA (Modifier or additive use); USES (Uses)
        (solid state lithium batteries based on organically
        modified ceramic electrolytes)
IT
     1871-21-2, ChlorotriVinylsilane
                                       2530-83-8,
     3-Glycidoxypropyltrimethoxysilane 4028-23-3, Allyl
                            7791-03-9, Lithium perchlorate 33454-82-9,
     dimethylchlorosilane
     Lithium trifluoromethanesulfonate
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (solid state lithium batteries based on organically
        modified ceramic electrolytes)
REFERENCE COUNT:
                               THERE ARE 7 CITED REFERENCES AVAILABLE FOR
```

THE RE FORMAT

THIS RECORD. ALL CITATIONS AVAILABLE IN

L27 ANSWER 19 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1998:13791 'HCAPLUS

DOCUMENT NUMBER:

128:91057

TITLE:

Electrodes for lithium-ion batteries

using polysiloxanes

INVENTOR(S):

Eguchi, Katsuya; Dahn, Jeffery Raymond; Wilson,

Alf M.; Xing, Weibing; Zank, Gregg Alan

PATENT ASSIGNEE(S):

Dow Corning Corporation, USA

SOURCE:

Eur. Pat. Appl., 19 pp.

DOCUMENT TYPE:

Patent

CODEN: EPXXDW

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 813258	A1	19971217	EP 1997-303911	199706
R: DE, FR, GB US 5824280	Α	19981020	US 1996-664278	06 199606
CA 2207290	AA	19971211	CA 1997-2207290	11 199706 09
JP 10074506	A2	19980317	JP 1997-153426	199706
PRIORITY APPLN. INFO.:			US 1996-664278	11 A 199606 11

AB The electrodes formed by the pyrolyzing a siloxane polymer (RR1R2SiO1/2)w(R3R4SiO)p(R5SiO3/2)q(SiO4/2)r at 700-1400°to form a ceramic material and by introducing Li ions into the formed material. R, R1, R2, R3, R4, and R5 are independently selected from H atom or C1-20 hydrocarbons, w is ≤ 0.8 , p is ≤ 0.9 , q is at ≤ 0.9 , r is ≤ 0.9 , and w + p + q + r = 1.

IT 1719-58-0DP, Chlorodimethylvinylsilane, reaction products with siloxanes

RL: PEP (Physical, engineering or chemical process); PNU

```
(Preparation, unclassified); PREP (Preparation); PROC (Process)
        (electrodes for lithium-ion batteries using pyrolyzed)
RN
     1719-58-0 HCAPLUS
CN
     Silane, chloroethenyldimethyl- (9CI) (CA INDEX NAME)
   Cl
Me-Si-CH-CH2
   Me
IC
     ICM H01M004-58
     52-2 (Electrochemical, Radiational, and Thermal Energy Technology)
CC
     Section cross-reference(s): 37, 57
     polysiloxane pyrolysis lithium ion battery electrode;
ST
     ceramics polysiloxane pyrolysis lithium battery electrode
IT
     Ceramics
        (electrodes for lithium-ion batteries using
        lithium-intercalatable pyrolyzed polysiloxane)
IT
     Battery anodes
        (lithium-intercalatable pyrolyzed polysiloxane)
IT
     Polysiloxanes, preparation
     RL: PEP (Physical, engineering or chemical process); PNU
     (Preparation, unclassified); PREP (Preparation); PROC (Process)
        (polysilalkylene-; electrodes for lithium-ion batteries
        using pyrolyzed)
ΙT
     Polycarbosilanes
     Polycarbosilanes
     RL: PEP (Physical, engineering or chemical process); PNU
     (Preparation, unclassified); PREP (Preparation); PROC (Process)
        (polysilalkylenes, siloxane-; electrodes for lithium-ion
        batteries using pyrolyzed)
ÌΤ
     Polysiloxanes, preparation
     RL: PEP (Physical, engineering or chemical process); PNU
     (Preparation, unclassified); PREP (Preparation); PROC (Process)
        (polysilphenylene-; electrodes for lithium-ion batteries
        using pyrolyzed)
     Polycarbosilanes
IT
     Polycarbosilanes
     RL: PEP (Physical, engineering or chemical process); PNU
     (Preparation, unclassified); PREP (Preparation); PROC (Process)
        (polysilphenylenes, siloxane-; electrodes for lithium-ion
```

batteries using pyrolyzed)

IT 1719-58-0DP, Chlorodimethylvinylsilane, reaction products 2627-95-4DP, 1,1,3,3-Tetramethyl-1,3with siloxanes divinyldisiloxane, reaction products with siloxanes 3277-26-7DP, 1,1,3,3-Tetramethyldisiloxane, reaction products with siloxanes 15545-80-9DP, 1,1,3,3-Tetraphenyldisiloxane, reaction products with 17306-05-7DP, reaction products with siloxanes siloxanes 18769-05-6DP, reaction products with siloxanes 25498-03-7DP, Methyltrimethoxysilane homopolymer, vinyl-terminated 29382-69-2DP. Vinyltrimethoxysilane homopolymer, vinyl-terminated 89885-26-7DP, Phenyltrimethoxysilane homopolymer, vinyl-terminated 162816-07-1DP, Methyltrimethoxysilane-phenyltrimethoxysilane 174305-53-4DP, trimethylsilylcopolymer, vinyl-terminated terminated 200959-33-7P 200959-35-9P 200959-36-0DP, vinyl-terminated RL: PEP (Physical, engineering or chemical process); PNU (Preparation, unclassified); PREP (Preparation); PROC (Process) (electrodes for lithium-ion batteries using pyrolyzed)

L27 ANSWER 20 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1997:743669 HCAPLUS

DOCUMENT NUMBER: 128:67667

TITLE: Cyclic voltammetric study of redox-active

surfactant by hydrogel-modified electrode

AUTHOR(S): Takeoka, Yukikazu; Aoki, Takashi; Sanui, Kohei;

Ogata, Naoya; Watanabe, Masayoshi

CORPORATE SOURCE: Department of Chemistry, Sophia University,

Tokyo, 102, Japan

SOURCE: Polymer Gels and Networks (1997), 5(4), 369-383

CODEN: PGNEEI; ISSN: 0966-7822

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

The dissolved states of redox-active non-ionic surfactant (FPEG) in the swollen state of N-iso-Pr acrylamide (NIPA) hydrogel were studied by using a gel-modified electrode. The pronounced decrease in the peak current and the neg. shift in the formal potential of cyclic voltammetry at the gel-modified electrode, as compared with the normal glassy carbon electrode, was obsd. in the micelle-soln.; this result indicates that the diffusive FPEG mols. which form the micelle hardly penetrate into the NIPA gel. This result suggests that there exists an interaction between FPEG mols. and the NIPA gel in the vicinity of the surface of the NIPA gel in the micelle-soln. However, this also indicates that a small amt. of FPEG mols. which can form micelles exists in the NIPA gel.

IT 1719-58-0, Chlorodimethylvinylsilane

RL: NUU (Other use, unclassified); PEP (Physical, engineering or chemical process); PRP (Properties); PROC (Process); USES (Uses) (silane coupler; in cyclic voltammetric study of redox-active surfactant by hydrogel-modified electrode)

RN 1719-58-0 HCAPLUS

CN Silane, chloroethenyldimethyl- (9CI) (CA INDEX NAME)

CC 72-2 (Electrochemistry)

IT 1719-58-0, Chlorodimethylvinylsilane

RL: NUU (Other use, unclassified); PEP (Physical, engineering or chemical process); PRP (Properties); PROC (Process); USES (Uses) (silane coupler; in cyclic voltammetric study of redox-active surfactant by hydrogel-modified electrode)

IT 7647-15-6, Sodium bromide, uses

RL: NUU (Other use, unclassified); PEP (Physical, engineering or chemical process); PROC (Process); USES (Uses)

(supporting electrolyte; cyclic voltammetric study of redox-active surfactant by hydrogel-modified electrode in soln. contg.)

REFERENCE COUNT:

15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L27 ANSWER 21 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1996:585618 HCAPLUS

DOCUMENT NUMBER:

125:275968

TITLE:

Electrochemical synthesis of

bis(2-thienyl)silanes, 2-thienylchlorosilanes,

bis(5-bromo-2-thienyl)silanes, and

(5-bromo-2-thienyl)dimethylchlorosilane, precursors of poly[(silanylene)thiophene]s Moreau, Carole; Serein-Spirau, Francoise; Bordeau, Michel; Biran, Claude; Dunogues,

Jacques

CORPORATE SOURCE:

Laboratoire Chimie Organique Organometallique,

Universite Bordeaux, Talance, F-33405, Fr.

SOURCE:

AUTHOR(S):

Journal of Organometallic Chemistry (1996),

522(2), 213-221

CODEN: JORCAI; ISSN: 0022-328X

PUBLISHER: Elsevier
DOCUMENT TYPE: Journal
LANGUAGE: English

OTHER SOURCE(S): CASREACT 125:275968

AB Bis(2-thienyl)silanes, e.g., bis(2-thienyl)dimethylsilane, and bis(5-bromo-2-thienyl)silanes were synthesized by electrochem. redn. of monohalothiophenes (Br, Cl) and 2,5-dibromothiophene resp. in the presence of a dichlorosilane in THF or DME, using an undivided cell, a sacrificial Mg or Al anode, a const. c.d. and Bu4NBr as the supporting electrolyte. When dichlorosilanes were used in large excess, halothiophenes underwent solely a monocoupling reaction leading selectivity to thienylchlorosilanes, which reveals the versatility of the electrochem. method.

IT 124-70-9, Dichloromethylvinylsilane

RL: RCT (Reactant); RACT (Reactant or reagent) (electrochem. redn. of halothiophenes in presence of chlorosilanes)

RN 124-70-9 HCAPLUS

CN Silane, dichloroethenylmethyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \texttt{C1} \\ | \\ \texttt{Me-Si-CH----} \texttt{CH}_2 \\ | \\ \texttt{C1} \end{array}$$

CC 29-6 (Organometallic and Organometalloidal Compounds)

Section cross-reference(s): 72

TT 75-78-5 80-10-4 96-43-5, 2-Chlorothiophene 124-70-9,
Dichloromethylvinylsilane 149-74-6 1003-09-4, 2-Bromothiophene
3141-27-3 7787-85-1, Dichloro(2-chloroethyl)methylsilane
13528-93-3, 1,2-Bis(chlorodimethylsilyl)ethane 18028-96-1
RL: RCT (Reactant); RACT (Reactant or reagent)
(electrochem. redn. of halothiophenes in presence of chlorosilanes)

L27 ANSWER 22 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:27541 HCAPLUS

DOCUMENT NUMBER: 124:158593

TITLE: Electrochemically induced silylation of

unsaturated compounds

AUTHOR(S):

Jouikov, V.; Grigorieva, L.

CORPORATE SOURCE:

Physical Chem. Dep., Kazan State Univ., Kazan,

420008, Russia

SOURCE:

Electrochimica Acta (1996), 41(3), 469-70

CODEN: ELCAAV; ISSN: 0013-4686

PUBLISHER: DOCUMENT TYPE: Elsevier Journal

LANGUAGE:

English

Electroredn. of some alkylchlorosilanes in the presence of difficult AB to reduce unsatd. compds. (phenylacetylene, styrene, cyclohexene) results in silylated products where the multiple bond has either reduced multiplicity (hydrosilylation) or the same multiplicity (silylation) depending on the nature of the starting compds. and conditions of electrolysis.

IT 124-70-9

> RL: NUU (Other use, unclassified); PRP (Properties); RCT (Reactant); RACT (Reactant or reagent); USES (Uses)

(in electrochem. silylation of styrene)

RN 124-70-9 HCAPLUS

CN Silane, dichloroethenylmethyl- (9CI) (CA INDEX NAME)

$$C1$$
 $|$
 $Me-Si-CH-CH_2$
 $|$
 $C1$

CC 72-2 (Electrochemistry)

Section cross-reference(s): 29

75-78-5, Dichlorodimethylsilane **124-70-9** 10026-04-7, IT

Silicon chloride sicl4

RL: NUU (Other use, unclassified); PRP (Properties); RCT (Reactant);

RACT (Reactant or reagent); USES (Uses)

(in electrochem. silylation of styrene)

L27 ANSWER 23 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1993:104052 HCAPLUS

DOCUMENT NUMBER:

118:104052

TITLE:

Molecular materials: a new type of polar

membrane for the titration of ions

AUTHOR(S):

Soulie, Corinne; Favier, Jean Claude; Hemery,

Patrick; Simon, Jacques

CORPORATE SOURCE:

CNRS, Paris, 75231, Fr.

SOURCE: Journal of Materials Chemistry (1992), 2(12),

1271-5

CODEN: JMACEP; ISSN: 0959-9428

DOCUMENT TYPE: Journal LANGUAGE: English

AB The synthesis of polar membranes having low glass transition temp. is described. They are obtained by room-temp. crosslinking of 2 cyanopropylsiloxane copolymers contg. .tplbond.SiCH:CH2 and .tplbond.SiH reactive groups. Conventional ionophores and electrolytes may be molecularly dissolved in the copolymers. The crosslinked membranes are used as a supporting matrix in ion-selective electrode devices. The best results are obtained when benzo-15-crown-5 is employed as ionophore.

IT 124-70-9

RL: RCT (Reactant); RACT (Reactant or reagent) (hydrolytic polymn. of, for prepn. of polar membranes for titrn. of ions)

RN 124-70-9 HCAPLUS

CN Silane, dichloroethenylmethyl- (9CI) (CA INDEX NAME)

CC 38-3 (Plastics Fabrication and Uses)

Section cross-reference(s): 72

IT 143-66-8

RL: USES (Uses)

(electrolyte, polar siloxane membranes contg., for titrn. of ions)

IT 75-54-7 **124-70-9** 1190-16-5

RL: RCT (Reactant); RACT (Reactant or reagent) (hydrolytic polymn. of, for prepn. of polar membranes for titrn. of ions)

L27 ANSWER 24 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1992:513684 HCAPLUS

DOCUMENT NUMBER: 117:113684

TITLE: Fluorocarbon-based polymer lamination coating

film and method of manufacturing the same

INVENTOR(S): Soga, Mamoru; Mino, Norihisa; Ogawa, Kazufumi;

PATENT ASSIGNEE(S):

Mochizuki, Yusuke; Shibata, Tsuneo

Matsushita Electric Industrial Co., Ltd., Japan

SOURCE:

Eur. Pat. Appl., 14 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
EP 484886				_	199111
EP 484886 R: DE, FR, GB,		19970312			05
US 5238746	A	19930824	US 1991-786824		199111 01
CA 2054931	AA	19920507	CA 1991-2054931		199111 05
CA 2054931	С		·		
JP 05064859	A2	19930319	JP 1991-288663		199111 05
JP 2528228 US 5587209		19960828 19961224	US 1995-438875		199505
DDIODIMU ADDINI TNEO	•		TD 1000 00001	_	10
PRIORITY APPLN. INFO.:			JP 1990-302021	A	199011 06
			US 1991-786824	А3	199111 01
				B1	199305 14
			US 1994-278126	В1	199407

21

AB The title method of applying an antifriction layer with good adhesion without electrolytic etching of substrates selected from metal, ceramic, glass, or plastic comprises first applying on the substrate a polymer absorption intermediate layer having Si bonds from dehydrochlorination or dealcoholation of silane coupling compd. and then laminating with a crosslinkable F-based polymer. An Al plate was dipped into a soln. of ω-nonadecenyltrichlorsilane in hexadecene-CCl4-CHCl3, washed, spray coated with [(CF2CF2)8CH:CH]n, and baked at 380° and irradiated with electron beam.

IT 125282-19-1

RL: USES (Uses)

(solns. of, for coating of aluminum before laminating with fluoropolymer layer)

RN 125282-19-1 HCAPLUS

CN Silane, trichloro-18-nonadecenyl- (9CI) (CA INDEX NAME)

 $H_2C = CH - (CH_2)_{17} - SiCl_3$

IC ICM B05D001-20

ICS C09D127-12

CC 42-10 (Coatings, Inks, and Related Products)

IT **125282-19-1** 143257-37-8

RL: USES (Uses)

(solns. of, for coating of aluminum before laminating with fluoropolymer layer)

L27 ANSWER 25 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1992:161234 HCAPLUS

DOCUMENT NUMBER: 116:161234

TITLE: Electrochemical manufacture of disilane

INVENTOR(S): Shono, Tatsuya; Kashiwamura, Shigefumi; Nishida,

Ryoichi

PATENT ASSIGNEE(S): Osaka Gas Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 9 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
			10011105		
	JP 03264683	A2	19911125	JP 1990-158224	199006 15
	JP 3016087	B2	20000306		
•	PRIORITY APPLN. INFO.:			JP 1990-9946 .A	1 199001 18

OTHER SOURCE(S): MARPAT 116:161234

AB A method for manufg. a disilane at a high yield involves a.c. or d.c. electrolysis of RR1R2SiX (R, R1, R2 = H, alkyl, aryl, alkoxy, amino; X = halo) using a Mg, Cu, Zn, Sn, or Al electrode(s). Addnl., ultrasound may be applied during the electrolysis.

IT 4028-23-3, Allyldimethylchlorosilane

RL: RCT (Reactant); RACT (Reactant or reagent)

(electrolysis of)

RN 4028-23-3 HCAPLUS

CN Silane, chlorodimethyl-2-propenyl- (9CI) (CA INDEX NAME)

IC ICM C25B003-02

ICS C07F007-08

CC 72-9 (Electrochemistry)
Section cross-reference(s): 49

ST disilane manuf electrolysis

Triphenylchlorosilane, reactions 76-86-8, Triphenylchlorosilane 768-33-2, Dimethylphenylchlorosilane 994-07-0, Methyldimethoxychlorosilane 1631-82-9, Phenylmethylchlorosilane 4028-23-3,

Allyldimethylchlorosilane 18162-84-0, Octyldimethylchlorosilane 139959-09-4

RL: RCT (Reactant); RACT (Reactant or reagent)
 (electrolysis of)

L27 ANSWER 26 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1992:135528 HCAPLUS

DOCUMENT NUMBER:

116:135528

TITLE:

Performance-oriented packaging standards;

changes to classification, hazard communication, packaging and handling requirements based on UN

standards and agency initiative

CORPORATE SOURCE:

United States Dept. of Transportation,

Washington, DC, 20590-0001, USA

SOURCE:

Federal Register (1990), 55(246), 52402-729, 21

Dec 1990

CODEN: FEREAC; ISSN: 0097-6326

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB The hazardous materials regulations under the Federal Hazardous Materials Transportation Act are revised based on the United Nations recommendations on the transport of dangerous goods. regulations cover the classification of materials, packaging requirements, and package marking, labeling, and shipping documentation, as well as transportation modes and handling, and incident reporting. Performance-oriented stds. are adopted for packaging for bulk and nonbulk transportation, and SI units of measurement generally replace US customary units. Hazardous material descriptions and proper shipping names are tabulated together with hazard class, identification nos., packing group, label required, special provisions, packaging authorizations, quantity limitations, and vessel stowage requirements.

75-94-5, Vinyltrichlorosilane 107-37-9, IT

Allyltrichlorosilane

RL: ADV (Adverse effect, including toxicity); PEP (Physical, engineering or chemical process); BIOL (Biological study); PROC (Process)

(packaging and transport of, stds. for)

75-94-5 HCAPLUS RN

CN Silane, trichloroethenyl- (9CI) (CA INDEX NAME)

 $Cl_3Si-CH=CH_2$

RN 107-37-9 HCAPLUS

CN Silane, trichloro-2-propenyl- (9CI) (CA INDEX NAME)

 $Cl_3Si-CH_2-CH \longrightarrow CH_2$

CC 59-6 (Air Pollution and Industrial Hygiene)

IT Adhesives

Alcoholic beverages

Ammunition

Antifreeze substances

Bactericides, Disinfectants, and Antiseptics

Batteries, primary

Blasting gelatin

Bombs (explosives)

Carbon paper

Cartridges

Castor bean

Coating materials

Corrosive substances

Cotton

Creosote

Detonators

Dyes

Dynamite

Electric fuses

Exothermic materials

Explosives

Flavoring materials

Flue dust

Fuel cells

Fuel oil

Fuels, diesel

Fuels, jet aircraft

Fusel oil

Fuses, explosives

Gas oils

Hay

Herbicides

Igniters and Lighters

Insecticides

Lacrimators

Magnetic substances

Matches

Oxidizing agents

Perfumes

Pesticides

Petroleum products

Pharmaceuticals

Photoelectric devices

Primers, explosive Projectiles Pyrophoric substances Pyrotechnic compositions Radioactive substances Refrigerating apparatus Rockets Shale oils Solvent naphtha Sprays Straw Textiles Thermoelectric devices Torpedoes (weapons) Turpentine Wood preservatives (packaging and transport of, stds. for) IT 50-00-0, Formaldehyde, miscellaneous 54-11-5, Nicotine 54-11-5D, Nicotine, compds. 55-63-0, Nitroglycerin 55-68-5, Phenylmercuric nitrate 56-18-8, 3,3'-Iminodipropylamine 56-23-5, miscellaneous 56-38-2, Parathion 57-06-7, Allyl isothiocyanate 57-14-7 57-24-9D, Strychnine, salts 60-00-4, EDTA, miscellaneous 60-29-7, Diethyl ether, miscellaneous 60-34-4, Methylhydrazine 60-57-1, Dieldrin 62-38-4, Phenylmercuric acetate 62-53-3, Aniline, miscellaneous 62-74-8, Sodium fluoroacetate 64-17-5, Ethanol, miscellaneous 64-18-6, Formic acid, miscellaneous 64-18-6D, Formic acid, chloro derivs. 64-19-7, Acetic acid, 64-67-5, Diethyl sulfate 66-25-1, Hexaldehyde miscellaneous 67-56-1, Methanol, miscellaneous 67-63-0, Isopropanol, miscellaneous 67-64-1, Acetone, miscellaneous 67-66-3, Chloroform, miscellaneous 68-11-1, Thioglycolic acid, miscellaneous 68-12-2, N,N-Dimethylformamide, miscellaneous 70-11-1, Phenacyl bromide 70-30-4, Hexachlorophene 71-23-8, n-Propanol, miscellaneous 71-41-0, 1-Pentanol, miscellaneous 71-43-2, Benzene, miscellaneous 71-55-6, 1,1,1-Trichloroethane 74-82-8, Methane, miscellaneous 74-83-9, miscellaneous 74-84-0, 74-85-1, Ethylene, miscellaneous Ethane, miscellaneous 74-86-2, Acetylene, miscellaneous 74-87-3, Methyl chloride, miscellaneous 74-88-4, Methyl iodide, miscellaneous 74-89-5, Methylamine, miscellaneous 74-90-8, Hydrogen cyanide, miscellaneous 74-93-1. Methyl mercaptan, miscellaneous 74-95-3, Dibromomethane 74-96-4, Ethyl bromide 74-97-5, Bromochloromethane 74-98-6, Propane, miscellaneous 75-00-3, Ethyl chloride 75-01-4, miscellaneous 75-02-5, Vinyl fluoride 75-04-7, Ethylamine, miscellaneous

Poisons

75-05-8, Methyl cyanide, miscellaneous 75-07-0, Acetaldehyde, miscellaneous 75-08-1, Ethyl mercaptan 75-09-2, Dichloromethane, miscellaneous 75-15-0, Carbon disulfide, miscellaneous 75-16-1, Methyl magnesium bromide 75-18-3, Dimethyl sulfide 75-19-4, 75-20-7, Calcium carbide Cyclopropane 75-21-8, Ethylene oxide, miscellaneous 75-21-8 75-25-2, Bromoform 75-26-3, 2-Bromopropane 75-28-5, Isobutane 75-28-5D, Isobutane, mixts. 75-29-6, 2-Chloropropane 75-31-0, Isopropylamine, miscellaneous 75-33-2, Isopropyl mercaptan 75-34-3, 1,1-Dichloroethane 75-35-4, miscellaneous 75-36-5, Acetyl chloride 75-38-7, 1,1-Difluoroethylene 75-39-8, Acetaldehyde ammonia 75-43-4. Dichloromonofluoromethane 75-44-5, Phosgene 75-45-6, Chlorodifluoromethane 75-46-7, Trifluoromethane 75-50-3, Trimethylamine, miscellaneous 75-52-5, Nitromethane, miscellaneous 75-54-7, Methyldichlorosilane 75-55-8, Propylenimine 75-56-9, Propylene oxide, miscellaneous 75-59-2, Tetramethylammonium 75-60-5, Cacodylic acid 75-61-6, hydroxide Dibromodifluoromethane 75-63-8 75-71-8, Dichlorodifluoromethane 75-72-9, Chlorotrifluoromethane 75-73-0, Tetrafluoromethane 75-76-3, Tetramethylsilane 75-77-4, Trimethylchlorosilane, miscellaneous 75-78-5, Dimethyldichlorosilane 75-79-6, Methyltrichlorosilane 75-83-2 75-86-5, Acetone cyanohydrin 75-87-6, Chloral 75-91-2, tert-Butyl hydroperoxide 75-94-5 , Vinyltrichlorosilane 76-01-7, Pentachloroethane Trichloroacetyl chloride 76-03-9, properties 76-05-1, Trifluoroacetic acid, miscellaneous 76-06-2, Chloropicrin 76-06-2D, Chloropicrin, mixts. 76-15-3 76-16-4, Hexafluoroethane 76-19-7, Octafluoropropane 76-22-2, Camphor 77-47-4, 77-73-6 77-78-1, Dimethyl sulfate Hexachlorocyclopentadiene 78-00-2, Tetraethyl lead 78-10-4, Tetraethyl silicate Dimethyldiethoxysilane 78-67-1, Azodiisobutyronitrile 2-Bromobutane 78-78-4, Isopentane 78-79-5, Isoprene, miscellaneous 78-81-9, Isobutylamine 78-82-0, Isobutyronitrile 78-83-1, Isobutanol, miscellaneous 78-84-2, Isobutyraldehyde 78-85-3, Methacrylaldehyde 78-87-5, Propylene dichloride 78-89-7, Propylene chlorohydrin 78-90-0, 1,2-Propylenediamine 78-93-3, 2-Butanone, miscellaneous 78-94-4, Methyl vinyl ketone, 78-95-5, Monochloroacetone miscellaneous 79-01-6, Trichloroethylene, miscellaneous 79-03-8, Propionyl chloride 79-04-9, Chloroacetyl chloride 79-06-1, Acrylamide, miscellaneous 79-08-3, Bromoacetic acid 79-09-4, Propionic acid, miscellaneous 79-10-7, 2-Propenoic acid, miscellaneous 79-11-8, Chloroacetic acid, miscellaneous 79-20-9, Methyl acetate 79-21-0, Peroxyacetic acid 79-22-1 79-24-3, Nitroethane 79-29-8, 2,3-Dimethylbutane 79-30-1, Isobutyryl chloride 79-31-2,

Isobutyric acid 79-36-7, Dichloroacetyl chloride 79-41-4, miscellaneous 79-42-5 79-43-6, Dichloroacetic acid, 79-44-7, Dimethylcarbamoyl chloride miscellaneous 80-10-4, Diphenyldichlorosilane 80-15-9, Cumene hydroperoxide Benzene sulfohydrazide 80-47-7, p-Menthane hydroperoxide 80-51-3, Diphenyloxide-4,4'-disulfohydrazide 80-56-8, 80-62-6 81-15-2 82-71-3 85-44-9, 86-50-0, Azinphos methyl 1,3-Isobenzofurandione 87-68-3, Hexachlorobutadiene 87-90-1 88-17-5, 2-Trifluoromethylaniline 88-72-2, o-Nitrotoluene 88-73-3, o-Chloronitrobenzene o-Nitroaniline 88-75-5, o-Nitrophenol 88-89-1 89-58-7, p-Nitroxylene 91-17-8, Decahydronaphthalene 91-20-3, Naphthalene, miscellaneous 91-20-3D, Naphthalene, diozonide 91-22-5, Quinoline, miscellaneous 91-59-8, β-Naphthylamine 91-66-7, N,N-Diethylaniline 92-52-4D, Biphenyl, chloro derivs. 92-52-4D, Biphenyl, halo derivs. 92-59-1, N-Ethyl-N-benzylaniline 92-87-5, Benzidine 93-58-3, Methyl benzoate 94-17-7, p-Chlorobenzoyl peroxide Benzoyl peroxide, miscellaneous 95-48-7, miscellaneous 95-50-1. o-Dichlorobenzene 95-54-5, o-Phenylenediamine, miscellaneous 95-55-6, o-Aminophenol 95-80-7 95-85-2, 2-Amino-4-chlorophenol 96-12-8, Dibromochloropropane 96-22-0, Diethyl ketone 96-23-1 96-24-2, Glycerol α-monochlorohydrin 96-32-2, Methyl bromoacetate 96-33-3 96-34-4, Methyl chloroacetate Methyl cyclopentane 96-41-3, Cyclopentanol 97-62-1. Ethyl isobutyrate 97-63-2 97-64-3, Ethyl lactate 97-72-3, Isobutyric 97-85-8, Isobutyl isobutyrate anhydride 97-86-9 97-88-1 97-96-1, 2-Ethylbutyraldehyde 98-00-0, Furfuryl alcohol 98-01-1, Furfural, miscellaneous 98-07-7, Benzotrichloride 98-08-8, Benzotrifluoride 98-09-9, Benzene sulfonyl chloride 98-12-4, Cyclohexyltrichlorosilane 98-13-5, Phenyltrichlorosilane 98-16-8, 3-Trifluoromethylaniline 98-82-8, Isopropylbenzene 98-83-9, miscellaneous 98-85-1, α -Methylbenzyl alcohol 98-87-3, Benzylidene chloride 98-88-4, Benzoyl chloride 98-95-3, Nitrobenzene, miscellaneous 99-08-1, m-Nitrotoluene 99-09-2, m-Nitroaniline 99-35-4, Trinitrobenzene 99-99-0, p-Nitrotoluene 100-00-5 100-01-6, p-Nitroaniline, miscellaneous 100-02-7, p-Nitrophenol, miscellaneous 100-17-4 100-34-5, Benzene diazonium chloride RL: ADV (Adverse effect, including toxicity); PEP (Physical, engineering or chemical process); BIOL (Biological study); PROC (Process) (packaging and transport of, stds. for) 100-36-7, N,N-Diethylethylenediamine 100-37-8, Diethylaminoethanol

100-39-0, Benzyl bromide 100-41-4, Ethylbenzene, miscellaneous

IT

100-42-5, miscellaneous 100-44-7, Benzyl chloride, miscellaneous 100-47-0, Benzonitrile, miscellaneous 100-50-5, 1,2,3,6-Tetrahydrobenzaldehyde 100-57-2, Phenylmercuric hydroxide 100-61-8, N-Methylaniline, miscellaneous 100-63-0, Phenylhydrazine 100-66-3, Anisole, miscellaneous 100-73-2, Acrolein dimer 101-25-7, N,N'-Dinitrosopentamethylenetetramine 101-68-8 101-77-9, 4,4'-Diaminodiphenyl methane 101-83-7, Dicyclohexylamine 102-69-2, Tripropylamine 102-70-5, Triallylamine 102-81-8, Dibutylaminoethanol 102-82-9, Tributylamine 103-65-1, 103-69-5, N-Ethylaniline n-Propylbenzene 103-71-9, Phenylisocyanate, miscellaneous 103-80-0, Phenylacetyl chloride 103-83-3, Benzyldimethylamine 104-15-4, Toluene sulfonic acid, miscellaneous 104-51-8, Butylbenzene 104-75-6, 2-Ethylhexylamine 104-78-9 104-90-5, 2-Methyl-5-ethylpyridine 105-36-2 105-37-3, Ethyl propionate 105-39-5, Ethyl chloroacetate 105-48-6, Isopropyl chloroacetate 105-54-4, Ethyl butyrate 105-56-6, Ethyl 105-57-7, Acetal cyanoacetate 105-58-8, Diethyl carbonate 105-64-6, Isopropyl peroxydicarbonate 105-74-8, Lauroyl peroxide 106-31-0, Butyric anhydride 106-44-5, p-Cresol, miscellaneous 106-46-7, p-Dichlorobenzene 106-50-3, p-Phenylenediamine, 106-51-4, 2,5-Cyclohexadiene-1,4-dione, miscellaneous miscellaneous 106-63-8, Isobutyl acrylate 106-68-3, Ethyl amyl 106-89-8, miscellaneous 106-88-7, 1,2-Butylene oxide 106-92-3, Allyl glycidyl ether 106-93-4, Ethylene dibromide 106-95-6, Allyl bromide, miscellaneous 106-96-7, 3-Bromopropyne 106-97-8, Butane, miscellaneous 106-97-8D, Butane, mixts. 106-99-0, 1,3-Butadiene, miscellaneous 107-00-6, Ethylacetylene 107-02-8, 2-Propenal, miscellaneous 107-05-1, Allyl chloride 107-06-2, Ethylene dichloride, miscellaneous 107-07-3, Ethylene chlorohydrin, miscellaneous 107-10-8, Propylamine, miscellaneous 107-11-9, Allylamine 107-12-0, Propionitrile 107-13-1, Acrylonitrile, miscellaneous 107-14-2, Chloroacetonitrile 107-15-3, Ethylenediamine, miscellaneous 107-18-6, Allyl alcohol, miscellaneous 107-19-7, Propargyl alcohol 107-20-0, 107-25-5, Vinylmethyl ether Chloroacetaldehyde 107-29-9, Acetaldehyde oxime 107-30-2, Methylchloromethyl ether Methyl formate 107-37-9, Allyltrichlorosilane 107-49-3, 107-71-1, tert-Butyl Tetraethyl pyrophosphate 107-70-0 peroxylacetate 107-72-2, Amyltrichlorosilane 107-81-3, 107-82-4, 1-Bromo-3-methylbutane 2-Bromopentane 107-87-9, Methyl propyl ketone 107-89-1, Aldol 107-92-6, Butyric acid, miscellaneous 108-01-0, Dimethylethanolamine 108-05-4, Acetic acid ethenyl ester, miscellaneous 108-09-8, 1,3-Dimethylbutylamine 108-10-1, Methyl isobutyl ketone 108-11-2, Methyl isobutyl carbinol 108-18-9, Diisopropylamine 108-20-3, Diisopropyl ether

108-21-4, Isopropyl acetate 108-22-5, Isopropenyl acetate 108-23-6, Isopropyl chloroformate 108-24-7, Acetic anhydride 108-31-6, 2,5-Furandione, miscellaneous 108-39-4, miscellaneous 108-45-2, m-Phenylenediamine, miscellaneous 108-46-3, Resorcinol. 108-67-8, miscellaneous miscellaneous 108-77-0 108-83-8, Diisobutyl ketone 108-84-9 108-86-1, Benzene, bromo-, miscellaneous 108-87-2, Methyl cyclohexane 108-88-3, Toluene, miscellaneous 108-90-7, Chlorobenzene, miscellaneous 108-91-8, 108-94-1, Cyclohexanone, Cyclohexylamine, miscellaneous 108-95-2, Phenol, miscellaneous miscellaneous 108-98-5, Phenyl mercaptan, miscellaneous 109-02-4 109-09-1, 2-Chloropyridine 109-13-7, tert-Butyl peroxyisobutyrate 109-52-4, Valeric acid, miscellaneous 109-53-5, Vinyl isobutyl ether 109-60-4, n-Propyl 109-61-5, n-Propyl chloroformate acetate 109-63-7, Boron trifluoride diethyl etherate 109-65-9, n-Butyl bromide 109-66-0, Pentane, miscellaneous 109-70-6, 1-Chloro-3-bromopropane 109-73-9, n-Butylamine, miscellaneous 109-74-0, Butyronitrile 109-77-3, Malononitrile 109-79-5, Butyl mercaptan 109-86-4, Ethylene glycol monomethyl ether 109-87-5, Methylal 109-89-7, Diethylamine, miscellaneous 109-90-0, Ethyl isocyanate 109-93-3, Divinyl ether 109-94-4, Ethyl Vinyl ethyl ether 109-95-5, Ethyl nitrite 109-99-9, Tetrahydrofuran, formate 110-00-9, Furan miscellaneous 110-01-0, Tetrahydrothiophene 110-02-1, Thiophene 110-12-3, 5-Methylhexan-2-one 110-16-7, Maleic acid, miscellaneous 110-18-9 110-19-0 110-22-5, 110-43-0, Amyl methyl ketone Diacetyl peroxide 110-49-6 110-54-3, Hexane, miscellaneous 110-58-7, Amylamine Valeraldehyde 110-66-7, Amyl mercaptan 110-68-9, 110-71-4, N-Methylbutylamine 110-69-0, Butyraldoxime 1,2-Dimethoxyethane 110-74-7, Propyl formate 110-78-1, n-Propyl 110-80-5, Ethylene glycol monoethyl ether 110-82-7, Cyclohexane, miscellaneous 110-83-8, Cyclohexene, miscellaneous 110-85-0, Piperazine, miscellaneous 110-86-1, Pyridine, 110-87-2 miscellaneous 110-89-4, Piperidine, miscellaneous 110-91-8, Morpholine, miscellaneous 110-96-3, Diisobutylamine 111-15-9, Ethylene glycol monoethyl ether acetate 111-34-2, Butylvinyl ether 111-36-4, n-Butyl isocyanate 111-40-0 111-49-9, Hexamethylenimine 111-43-3, Dipropyl ether 111-65-9, Octane, miscellaneous 111-69-3, Adiponitrile 111-71-7, n-Heptaldehyde 111-76-2, Ethylene glycol monobutyl ether 111-92-2, Di-n-butylamine 112-04-9 112-24-3, Triethylenetetramine 112-57-2 115-07-1, Propylene, miscellaneous 115-10-6, Dimethyl ether 115-11-7, Isobutylene, miscellaneous 115-21-9, Ethyltrichlorosilane 115-25-3, Octafluorocyclobutane 116-14-3, Tetrafluoroethylene, miscellaneous 116-15-4,

116-16-5, Hexachloroacetone 116-54-1, Methyl Hexafluoropropylene dichloroacetate 118-74-1, Hexachlorobenzene 118-96-7, 120-92-3, Cyclopentanone 121-43-7, Trimethyl Trinitrotoluene 121-44-8, Triethylamine, miscellaneous borate 121-45-9, 121-46-0, 2,5-Norbornadiene Trimethyl phosphite 121-69-7, N, N-Dimethylaniline, miscellaneous 121-73-3 121-82-4, Cyclotrimethylenetrinitramine 122-51-0, Ethyl orthoformate 122-52-1, Triethyl phosphite 123-00-2, 4-Morpholinepropanamine 123-19-3, Dipropylketone 123-20-6, Vinyl butyrate 123-23-9, Succinic acid peroxide 123-30-8, p-Aminophenol 123-31-9, Hydroquinone, miscellaneous 123-38-6, Propionaldehyde, miscellaneous 123-42-2, Diacetone alcohol 123-54-6, 2,4-Pentanedione, miscellaneous 123-62-6, Propionic anhydride 123-63-7, Paraldehyde 123-72-8, Butyraldehyde 123-75-1, Pyrrolidine, miscellaneous 123-86-4, Butyl acetate 123-91-1, Dioxane, miscellaneous 124-02-7, Diallylamine 124-09-4, Hexamethylenediamine, miscellaneous 124-13-0, Octyl aldehyde 124-18-5, n-Decane 124-38-9, Carbon dioxide, miscellaneous 124-40-3, Dimethylamine, miscellaneous 124-41-4, Sodium methylate 124-47-0, Urea nitrate 124-65-2, Sodium cacodylate 126-98-7, Methacrylonitrile 126-99-8, Chloroprene 127-18-4, Tetrachloroethylene, miscellaneous 127-85-5, Sodium arsanilate 131-52-2, Sodium pentachlorophenate 131-73-7, Hexanitrodiphenylamine 131-74-8, Ammonium picrate 133-14-2 133-55-1, N,N'-Dinitroso-N,N'-dimethyl terephthalamide α-Naphthylamine RL: ADV (Adverse effect, including toxicity); PEP (Physical, engineering or chemical process); BIOL (Biological study); PROC (Process)

(packaging and transport of, stds. for)

L27 ANSWER 27 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1988:28552 HCAPLUS

DOCUMENT NUMBER: 108:28552

TITLE: Modifying steel surfaces with alkoxy- and

chlorosilanes

AUTHOR(S): Nazarov, A. P.; Petrunin, M. A.; Mikhailovskii,

Yu. N.

CORPORATE SOURCE: Inst. Fiz. Khim., Moscow, USSR

SOURCE: Zashchita Metallov (1987), 23(6), 1007-11

CODEN: ZAMEA9; ISSN: 0044-1856

DOCUMENT TYPE: Journal LANGUAGE: Russian

AB An attempt was made to clarify the conditions of formation of chemisorption bonds between the surface atoms of the metal and the

mols. of silanes. The electrochem. properties were studied as modified by several different methods. The electrochem. method of study (plotting of polarization curves) is sensitive and informative concerning the formation of adsorption complexes or their assocs. The investigation was made on rotating disk electrodes of steel St.3 in the presence of the following silanes: Si(OEt)4, H2C:CHSiCl3, MeSiCl3, Me2SiCl2, and Me3SiOEt.

IT 75-94-5, Vinyltrichlorosilane

RL: PRP (Properties)

(anodic polarization of steel electrode modified with)

RN 75-94-5 HCAPLUS

CN Silane, trichloroethenyl- (9CI) (CA INDEX NAME)

 $Cl_3Si-CH=CH_2$

CC 72-11 (Electrochemistry)

Section cross-reference(s): 29, 55

IT **Electrolytic** polarization

(of steel surface-modified with silanes)

IT 75-78-5, Dimethyldichlorosilane 75-79-6, Methyltrichlorosilane 75-94-5, Vinyltrichlorosilane 78-10-4, Tetraethoxysilane

1825-62-3, Trimethylethoxysilane

RL: PRP (Properties)

(anodic polarization of steel electrode modified with)

IT 7757-82-6, Disodium sulfate, properties

RL: PRP (Properties)

(electrolytic polarization of silane-modified steel electrodes in soln. contg.)

L27 ANSWER 28 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1965:413469 HCAPLUS

DOCUMENT NUMBER: 63:13469
ORIGINAL REFERENCE NO.: 63:2391e-g

TITLE: Determination of double bonds in organosilicon

compounds, containing SiH groups

AUTHOR(S): Kreshkov, A. P.; Bork, V. A.; Aparsheva, M. I.

SOURCE: Plasticheskie Massy (1965), (4), 63-5

CODEN: PLMSAI; ISSN: 0554-2901

DOCUMENT TYPE: Journal LANGUAGE: Russian

AB A rapid accurate, and sufficiently sensitive method is described for detn. of double bonds to 10-3% in organo-Si compds. contg. SiH groups. The SiH bond was quant. destroyed by KOH-MeOH soln. and

then the double bonds were titrated amperometrically with Br soln. in MeOH. The SiH groups may be detd. by amperometric titrn. of the total amt. of SiH groups and double bonds without the previous The titrn. was carried out at 0.3 v. with 4M HCl as destruction. supporting electrolyte. A rotating Pt microelectrode (900 rpm.) was used as anode; a S.C.E. served as cathode. The relative error was 2%. Two 10-mg. sample was dissolved in 3 ml. of concd. HCl and the soln. was dild. to 25 ml. with 0.05M NaBr in MeOH. soln. was carefully mixed and transferred into the polarographic cell; 0.1M soln. of Br in MeOH, satd. with NaBr, was used as titrn. reagent. After the addn. of each portion of titrn. reagent the soln. was mixed and after 30-40 sec. the height of the polarographic wave was registered. To 10-30 mg. sample in MeOH was added 5 ml. 2N KOH in MeOH; after 3 hrs., the soln. was dild. to 25 ml. with MeOH. Three to 5 ml. of this soln. were transferred into the polarographic cell, 3 ml. of concd. HCl were added, and the soln. was dild. to 20-5 ml. with MeOH. The height of the polarographic wave was recorded after 25 sec.

IT 124-70-9, Silane, dichloromethylvinyl-

(detn. of double bonds and SiH groups in mixts. of other organo-Si compds. and)

RN 124-70-9 HCAPLUS

CN Silane, dichloroethenylmethyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \texttt{C1} \\ | \\ \texttt{Me-Si-CH----} \texttt{CH2} \\ | \\ \texttt{C1} \end{array}$$

CC 2 (Analytical Chemistry)

IT 124-70-9, Silane, dichloromethylvinyl- 762-72-1, Silane, allyltrimethyl-

(detn. of double bonds and SiH groups in mixts. of other organo-Si compds. and)

L27 ANSWER 29 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1956:53894 HCAPLUS

DOCUMENT NUMBER: 50:53894

ORIGINAL REFERENCE NO.: 50:10309g-i,10310a

TITLE: Ion-exchange resin compositions

INVENTOR(S): Berry, Kenneth L.; Caddell, Jack R.

PATENT ASSIGNEE(S): E. I. du Pont de Nemours & Co.

DOCUMENT TYPE:

Patent

LANGUAGE:

Unavailable

FAMILY ACC. NUM. COUNT:

. 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE

US 2739906

19560327 US

A particulate, ion-exchange resin (I)-coated, inert solid with a sp. AB gr. ≥ 1.5 is composed of particles of an inert, nonporous, water-insol. inorg. solid having dimensions of 0.01-0.5 cm., a wt. \geq 75 lb./cu. ft., and a sp. surface \leq 250 sq. cm./q. These particles are at least partially coated with I. Thus, glass beads, 0.65 cm. in diam., were wet with vinyltrichlorosilane and the excess evapd. Treated beads (38 parts) were placed in a mixt. of 37 parts styrene, 1 part of 40% divinylbenzene in ethylvinylbenzene, and 0.25 part α,α' -azobis $(\alpha,\alpha,\alpha$ trimethylvaleronitrile), which was heated for about 10 hrs. at 35° under N until the liquid component became viscous and appeared to be at the point of incipient gelation. α, α' -Azobis (a, a-dimethylvaleronitrile) (0.1 part) and the mixt. are placed in a soln. of 10 parts methylcellulose ("Methocel," 4000 centipoises) in 400 parts H2O, which had been deaerated and blanketed by N. This mixt. (II) was agitated until the beads and the org. phase became dispersed. II was then heated for 4 hrs. at 60° and 2 hrs. at 100° to complete the polymerization to the infusible stage. II was dild. with H2O, and the polymer-coated beads (III) were recovered by washing with H2O, dried, and screened (8-mesh) to remove aggregates. III were treated in boiling H2O for 1 hr. to remove a trace of styrene. dried and placed in C2H4Cl2 (d. = 1.26), and the material that floated was discarded. The product was next dried and placed in MeI (d. = 2.28), and the material which sank was discarded. III were baked overnight in vacuo at 86°, placed in 95% H2SO4 for 60 hrs. at room temp., and then heated 15 min. in the acid at 110-20°. This step converted the polymer on the glass beads to I by sulfonation.

- IT 75-94-5, Silane, trichlorovinyl-
 - (glass beads treated with, for coating with ion-exchange resins)
- RN 75-94-5 HCAPLUS
- CN Silane, trichloroethenyl- (9CI) (CA INDEX NAME)

 $Cl_3Si-CH=CH_2$

- CC 13 (Chemical Industry and Miscellaneous Industrial Products)
- IT Ions

(electrolytic, -exchanging substances, glass beads coated with resinous)

=> d 128 ti 1-YOU HAVE REQUESTED DATA FROM 27 ANSWERS - CONTINUE? Y/(N):y

- L28 ANSWER 1 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Titanocene and zirconocene complexes containing dendrimersubstituted cyclopentadienyl ligands - synthesis and ethylene polymerization
- L28 ANSWER 2 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Metal oxide composites containing acrylate-based monomers
- L28 ANSWER 3 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Stepwise and dissociative mechanisms of the electron transfer in electrochemical reactions involving organosilicon compounds: molecular-thermodynamic approach
- L28 ANSWER 4 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Emulsions of fragrance releasing silicon compounds
- L28 ANSWER 5 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Comparison of methods for determination of acidic surface oxides on carbon blacks
- L28 ANSWER 6 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI 1,1'-Bis(dimethylvinylsilyl)ferrocene as a Two-Directional Core for the Construction of Homo- and Heterometallic Systems
- L28 ANSWER 7 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Selective and high-yield electrosynthesis of (silyl and silanylene 1-methylpyrroles) from 1-methylpyrrole bromides
- L28 ANSWER 8 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Dendrimers Containing Organometallic Moieties Electronically

Communicated

- L28 ANSWER 9 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Competitive electrochemical thio- and selenenylation of chlorosilanes
- L28 ANSWER 10 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Electrochemical synthesis of cyclic alkylsilanes
- L28 ANSWER 11 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Blueprint for a lipase support: use of hydrophobic controlled-pore glasses as model systems
- L28 ANSWER 12 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Manufacture of chemically adsorbed ultrathin fluorosiloxane films
- L28 ANSWER 13 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Studies of molecular alignments of monolayers deposited by a chemical adsorption technique
- L28 ANSWER 14 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Optical recording material
- L28 ANSWER 15 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Preparation of multilayers using chemical adsorption and electron-beam irradiation
- L28 ANSWER 16 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Reactions of chemically adsorbed monolayers induced by electron beam irradiation in active gas atmosphere and applications for the preparation of multilayers
- L28 ANSWER 17 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Producing a polyacetylene
- L28 ANSWER 18 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Mechanism of the grafing of organosilanes on mineral surfaces. II Secondary reaction during the grafting of alkenylchlorosilanes
- L28 ANSWER 19 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Silica antifoaming agents
- L28 ANSWER 20 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Methylvinylcyclotrisiloxane

- L28 ANSWER 21 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Waterproofing of materials by organosilicon compounds. XIV. Chemical stability of waterproof organosilicon films on glass
- L28 ANSWER 22 OF 27 HCAPLUS COPYRIGHT, 2006 ACS on STN
- TI Silicone waterproofing coatings
- L28 ANSWER 23 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Exterior paints
- L28 ANSWER 24 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Silicones. I. LXXIX. Preparation of organopentafluorosilicates in nonaqueous solvents
- L28 ANSWER 25 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Glass-fiber coatings
- L28 ANSWER 26 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Insulating electrical conductors
- L28 ANSWER 27 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN
- TI Coating compositions containing a copolymer of a vinyl polysiloxane, an unsaturated polyester resin, and a vinyl compound
- => d 128 ibib abs hitstr hitind 24

L28 ANSWER 24 OF 27 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1965:66646 HCAPLUS

DOCUMENT NUMBER: 62:66646

ORIGINAL REFERENCE NO.: 62:11844f-h,11845a-e

TITLE: Silicones. I. LXXIX. Preparation of

organopentafluorosilicates in nonaqueous

solvents

AUTHOR(S): Mueller, Richard; Dathe, Christian; Mross,

Dieter

CORPORATE SOURCE: Inst. Silikon-Fluorokarbon-Chem., Radebeul,

Dresden, Germany

SOURCE: Chemische Berichte (1965), 98(1), 241-4

CODEN: CHBEAM; ISSN: 0009-2940

DOCUMENT TYPE: Journal LANGUAGE: German

AB The new organopentafluorosilicates, such as (NH4)2[MeSiF5] (I) and K2[PhSiF5] (II), were prepd. without solvents, in H2O, and also in org. solvents. MeSiCl3 (300 g.) added dropwise during 24 h. to 400

g. SbF3 and heated 2 h. at 70-80° yielded 145 g. MeSiF3 (III), b. -29.7 to -29.2°. PhSiCl3 (422 g.) and 680 g. SbF3 refluxed 1 h. gave 85% PhSiF3 (IV), b. 102-3°, d20 1.210. III passed 8 h. into 20 q. NH4F in 250 cc. Me2CO, and the ppt. in H2O treated with nearly satd. aq. KF yielded K2 [MeSiF5]. Similarly were performed the following runs (listed in the table). , fluoride used, millimoles used, solvent used, cc., mg.-% H2O in solvent, reaction time (hrs.), III (g.)consumed, % reaction; NH4F, 270, MeCN, 100, 12.52, 16, 11, 82; NH4F, 270, C6H6, 100, 79.5, 8, 5, 37; NH4F, 270, petr.ether, 100, 48.4, 8, 2, 15; KF, 172, Me2CO, 120, 813, 8, 5, 58; KF, 344, MeCN, 150, 17.6, 8, 10, 58; KF, 172, C6H6, 150, 79.5, 8, 2, 22; KF, 86, petr. ether, 100, 48.4, 16, 0.25, 6; NaF, 119, MeCN, 100, 21.6, 6, 5.4, 90; NH4F (6 q.) in 100 cc. C6H6 contq. 79.5 mg. H2O/100 cc. treated dropwise with 14 g. IV and stirred 8 h. gave 9 g. mixt. of NH4F and (NH4) 2 [PhSiF5]. Similarly were performed the runs listed in the 2nd table. fluoride used, milli, millimoles used, millimoles IV used, solvent used, cc., mg.-% H2O in solvent, reaction time (hrs.), IV (g.) used, % reaction; NH4F, 325, 164, Me2CO, 100, 607, 3, 26, 99; NH4F, 325, 164, MeCN, 100, 17.6, 4, 25, 95; NH4F, 162, 87, petr. ether, 100, 48.4, 8, 3, 36; KF, 172, 87, MeCN, 100, 607, 5, 14, 100; KF, 172, 87, MeCN, 100, 12.52, 5, 12, 86; KF, 172, 87, C6H6, 100, 18.78, 5, 1, 10; KF, 172, 87, petr. ether, 110, 48.8, 5, 1, 11;

RN 16924-21-3 HCAPLUS

CN Silicate(2-), ethenylpentafluoro-, diammonium, (OC-6-21)- (9CI) (CA INDEX NAME)

●2 NH4+

39 (Organometallic and Organometalloidal Compounds) CC 368-47-8, Silane, trifluorophenyl- 373-74-0, Silane, IT trifluoromethyl- 730-93-8, Phosphonimidic diamide, N, N', N''-triisopropyl-P-phenyl- 884-76-4, Phosphonimidic diamide, N, N', N''-triethyl-P-phenyl- 1840-61-5, Ammonium pentafluoromethylsilicate, (NH4)2[SiF5-Me] 5507-59-5, Potassium pentafluorophenylsilicate 12105-98-5, Ammonium pentafluorophenylsilicate, (NH4)2[SiF5Ph] 12105-98-5, Ammonium pentafluorophenylsilicate, (NH4)2[SiF5Ph] 16924-21-3, Silicate, pentafluorovinyl-, diammonium 17979-70-3, Potassium pentafluoromethylsilicate, K2[SiF5-Me] 46059-39-6, Silicate, pentafluorophenyl-(prepn. of)